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ABSTRACTS OF THE 190TH AMERICAN CHEMICAL SOCIETY NATIONAL MEETING, vol. 190,1985, page 23, no. 47; R.R. BOTT et al.: "Protein engineering of subtilisin"

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JOURNAL OF CELLULAR BIOCHEMISTRY SUPPL., vol. 0, no. 10, part A, 1986, page271, no. E101, SYMPOSIUM ON PROTEASES IN BIOLOGICAL CONTROL AND BIOTECHNOLOGY,15th ANNUAL UCLA, MEETING ON MOLECULAR AND CELLULAR BIOLOGY, Los Angeles, CA.,9th-15th February 1986; P. BRY-AN et al.: "Protein engineering of subtilisin-proteases of enhanced stability"

WORLD BIOTECH. REPORT, vol. 2, 1985, pages 51-59, Online Publications, Pinner,GB; R. BOTT: "Modeling & crystallographic analysis of site-specific mutants of subtillsin"

JOURNAL OF CELLULAR BIOCHEMISTRY SUPPL., vol. 0, no. 11, part C, 1987, page 200, no. N024, New York, US; D.A. ESTELL et al.: "Tailoring enzymatic properties through multiple mutations"

PROCEEDINGS OF THE NATIONAL ACADEMY OF SCIENCE USA, vol. 84, March 1987, pages 1219-1223, Washington, D.C., US; J.A. WELLS et al.: "Designing substrate specifity by protein engineering of electrostatic interactions"

BIOCHEMISTRY, vol. 26, no. 8, April 1987, pages 2077-2082, American Chemical Society, Washington, D.C., US; M.W. PANTOLIANO et al.: "Protein engineering of subtilisin BPN': enhanced stabilization through the introduction of two cysteines to form a disulfide bond"

PROCEEDINGS OF THE NATIONAL ACADEMY OF SCIENCE USA, vol. 83, June 1986, pages 3743-3745, Washington, D.C., US; P. BRYAN et al.: "Site-directed mutagenesis and the role of the oxyanion hole in subtilisin"

NATURE, vol. 318, 28th November 1985, pages 375-376, London, GB; P.G. THOMAS etal.: "Tailoring the pH dependence of enzyme catalysis using proteinengineering"

JOURNAL OF BACTERIOLOGY, vol. 158, no. 2, May 1984, pages 411-418, American Society for Microbiology, Washington, D.C., US; M.L. STAHL et al.: "Replacement of the Bacillus subtilis subtilisin structural gene with an in vitro-derived deletion mutation"

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Representative: Armitage, Ian Michael et al MEWBURN ELLIS York House 23 Kingsway London WC2B 6HP (GB) NUCLEIC ACIDS RESEARCH, vol. 11, no. 22, November 1983, pages 7911-7925, IRL Press Ltd, Cambridge, GB; J.A. WELLS et al.: "Cloning, sequencing, and secretion of Bacillus amyloliquefaciens subtilisin in Bacillus subtilis"

Description

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The recent development of various in vitro techniques to manipulate the DNA sequences encoding naturally-occuring polypeptides as well as recent developments in the chemical synthesis of relatively short sequences of single and double stranded DNA has resulted in the speculation that such techniques can be used to modify enzymes to improve some functional property in a predictable way. Ulmer, K.M. (1983) Science 219, 666-671. The only working example disclosed therein is the substitution of a single amino acid within the active site of tyrosyl-tRNA synthetase (Cys35→Ser) which lead to a reduction in enzymatic activity. See Winter, G., et al. (1982) Nature 299, 756-758; and Wilkinson, A.J., et al. (1983) Biochemistry 22, 3581-3586 (Cys35→Gly mutation also resulted in decreased activity).

When the same t-RNA synthetase was modified by substituting a different amino acid residue within the active site with two different amino acids, one of the mutants (Thr51→Ala) reportedly demonstrated a predicted moderate increase in kcat/Km whereas a second mutant (Thr51→Pro) demonstrated a massive increase in kcat/Km which could not be explained with certainty. Wilkinson, A.H., et al. (1984) Nature 307, 187-188.

Another reported example of a single substitution of an amino acid residue is the substitution of cysteine for isoleucine at the third residue of T4 lysozyme. Perry, L.J., et al. (1984) Science 226, 555-557. The resultant mutant lysozyme was mildly oxidized to form a disulfide bond between the new cysteine residue at position 3 and the native cysteine at position 97. This crosslinked mutant was initially described by the author as being enzymatically identical to, but more thermally stable than, the wild type enzyme. However, in a "Note Added in Proof", the author indicated that the enhanced stability observed was probably due to a chemical modification of cysteine at residue 54 since the mutant lysozyme with a free thiol at Cys54 has a thermal stability identical to the wild type lysozyme.

Similarly, a modified dihydrofolate reductase from E.coli has been reported to be modified by similar methods to introduce a cysteine which could be cross linked with a naturally-occurring cysteine in the reductase. Villafranca, D.E., et al. (1983) Science 222, 782-788. The author indicates that this mutant is fully reactive in the reduced state but has significantly diminished activity in the oxidized state. In addition, two other substitutions of specific amino acid residues are reported which resulted in mutants which had diminished or no activity.

EPO Publication No. 0130756 discloses the substitution of specific residues within \underline{B} . $\underline{amyloliquefaciens}$ subtilisin with specific amino acids. Thus, Met222 has been substituted with all 19 other amino acids, Gly166 with 9 different amino acids and Gly169 with Ala and Ser.

As set forth below, several laboratories have also reported the use of site directed mutagensis to produce the mutation of more than one amino acid residue within a polypeptide.

The amino-terminal region of the signal peptide of the prolipoprotein of the <u>E. coli</u> outer membrane was stated to be altered by the substitution or deletion of residues 2 and 3 to produce a charge change in that region of the polypeptide. Inoyye, S., et al. (1982) <u>Proc. Nat. Acad. Sci. USA 79</u>, 3438-3441. The same laboratory also reported the substitution and deletion of amino acid redisues 9 and 14 to determine the effects of such substitution on the hydrophobic region of the same signal sequence. Inouye, S., et al. (1984) J. Biol. Chem. 259, 3729-3733.

Double mutants in the active site of tyrosyl-t-RNA synthetase have also been reported. Carter, P.J., et al. (1984) Cell 38, 835-840. In this report, the improved affinity of the previously described Thr51-Pro mutant for ATP was probed by producing a second mutation in the active site of the enzyme. One of the double mutants, Gly35/Pro51, reportedly demonstrated an unexpected result in that it bound ATP in the transition state better than was expected from the two single mutants. Moreover, the author warns, at least for one double mutant, that it is not readily predictable how one substitution alters the effect caused by the other substitution and that care must be taken in interpreting such substitutions.

A mutant is disclosed in U.S. Patent No. 4,532,207, wherein a polyarginine tail was attached to the C-terminal residue of \$\beta\$-urogastrone by modifying the DNA sequence encoding the polypeptide. As disclosed, the polyarginine tail changed the electrophoretic mobility of the urogastrone-polyaginine hybrid permiting selective purification. The polyarginine was subsequently removed, according to the patentee, by a polyarginine specific exopeptidase to produce the purified urogastrone. Properly construed, this reference discloses hybrid polypeptides which do not constitute mutant polypeptides containing the substitution, insertion or deletion of one or more amino acids of a naturally occurring polypeptide.

Single and double mutants of rat pancreatic trypsin have also been reported. Craik, C.S., et al. (1985) Science 228, 291-297. As reported, glycine residues at positions 216 and 226 were replaced with alanine residues to produce three trypsin mutants (two single mutants and one double mutant). In the case of the single mutants, the authors stated expectation was to observe a differential effect on Km. They instead

reported a change in specificity (kcat/Km) which was primarily the result of a decrease in kcat. In contrast, the double mutant reportedly demonstrated a differential increase in Km for lysyl and arginyl substrates as compared to wild type trypsin but had virtually no catalytic activity.

The references discussed above are provided solely for their disclosure prior to the filing date of the instant case, and nothing herein is to be construed as an admission that the inventors are not entitled to antedate such disclosure by virtue of prior invention or priority based on earlier filed applications.

Based on the above references, however, it is apparent that the modification of the amino acid sequence of wild type enzymes often results in the decrease or destruction of biological activity.

Accordingly, it is an object herein to provide carbonyl hydrolase mutants which have at least one property which is different from the same property of the carbonyl hydrolase precursor from which the amino acid of said mutant is derived.

It is a further object to provide mutant DNA sequences encoding such carbonyl hydrolase mutants as well as expression vectors containing such mutant DNA sequences.

Still further, another object of the present invention is to provide host cells transformed with such vectors as well as host cells which are capable of expressing such mutants either intracellularly or extracellularly.

Summary of the Invention

The invention includes carbonyl hydrolase mutants, preferably having at least one property which is substantially different from the same property of the precursor non-human carbonyl hydrolase from which the amino acid sequence of the mutant is derived. These properties include oxidative stability, substrate, specificity catalytic activity, thermal stability, alkaline stability, pH activity profile and resistance to proteolytic degradation. The precursor carbonyl hydrolase may be naturally occurring carbonyl hydrolases or recombinant carbonyl hydrolases. The amino acid sequence of the carbonyl hydrolase mutant is derived by the substitution, deletion or insertion of one or more amino acids of the precursor carbonyl hydrolase amino acid sequence.

The invention also includes mutant DNA sequences encoding such carbonyl hydrolase mutants. Further the invention includes expression vectors containing such mutant DNA sequences as well as host cells transformed with such vectors which are capable of expressing said carbonyl hydrolase mutants.

Brief Description of the Drawings

Figure 1 shows the nucleotide sequence of the coding strand, correlated with the amino acid sequence of B. amyloliquefaciens subtilisin gene. Promoter (p) ribosome binding site (rbs) and termination (term) regions of the DNA sequence as well as sequences encoding the presequence (PRE) putative prosequence (PRO) and mature form (MAT) of the hydrolase are also shown.

Figure 2 is a schematic diagram showing the substrate binding cleft of subtilisin together with substrate. Figure 3 is a stereo view of the S-1 binding subsite of B. amyloliquefaciens subtilisin showing a lysine P-1 substrate bound in the site in two different ways. Figure 3A shows Lysine P-1 substrate bound to form a salt bridge with a Glu at position 156. Figure 3B shows Lysine P-1 substrate bound to form a salt bridge with Glu at position 166.

Figure 4 is a schematic diagram of the active site of subtilisin Asp32, His64 and Ser221.

Figures 5A and 5B depict the amino acid sequence of subtilisin obtained from various sources. The residues directly beneath each residue of B. amyloliquefaciens subtilisin are equivalent residues which (1) can be mutated in a similar manner to that described for B. amyloliquefaciens subtilisin, or (2) can be used as a replacement amino acid residue in B. amyloliquefaciens subtilisin. Figure 5C depicts conserved residues of B. amyloliquefaciens subtilisin when compared to other subtilisin sequences.

Figures 6A and 6B depict the inactivation of the mutants Met222L and Met222Q when exposed to various organic oxidants.

Figure 7 depicts the ultraviolet spectrum of Met222F subtilisin and the difference spectrum generated after inactivation by diperdodecanoic acid (DPDA).

Figure 8 shows the pattern of cyanogen bromide digests of untreated and DPDA oxidized subtilisin Met222F on high resolution SDS-pyridine peptide gels.

Figure 9 depicts a map of the cyanogen bromide fragments of Fig. 8 and their alignment with the sequence of subtilisin Met222F.

Figure 10 depicts the construction of mutations between codons 45 and 50 of <u>B</u>. <u>amyloliquefaciens</u> subtilisin.

Figure 11 depicts the construction of mutations between codons 122 and 127 of <u>B</u>. <u>amyloliquefaciens</u> subtilisin.

Figure 12 depicts the effect of DPDA on the activity of subtilisin mutants at positions 50 and 124 in subtilisin Met222F.

Figure 13 depicts the construction of mutations at codon 166 of B. amyloliquefaciens subtilisin.

Figure 14 depicts the effect of hydrophobicity of the P-1 substrate side-chain on the kinetic parameters of wild-type B. amyloliquefaciens subtilisin.

Figure 15 depicts the effect of position 166 side-chain substitutions on P-I substrate specificity. Figure 15A shows position 166 mutant subtilisins containing non-branched alkyl and aromatic side-chain substitutions arranged in order of increasing molecular volume. Figure 15B shows a series of mutant enzymes progressing through β- and γ-branched aliphatic side chain substitutions of increasing molecular volume.

Figure 16 depicts the effect of position 166 side-chain volumn on log kcat/Km for various P-1 substrates.

Figure 17 shows the substrate specificity differences between Ile166 and wild-type (Gly166) B. amyloliquefaciens subtilisin against a series of alphatic and aromatic substrates. Each bar represents the difference in log kcat/Km for Ile166 minus wild-type (Gly166) subtilisin.

Figure 18 depicts the construction of mutations at codon 169 of B. amyloliquefaciens subtilisin.

Figure 19 depicts the construction of mutations at codon 104 of B. amyloliquefaciens subtilisin.

Figure 20 depicts the construction of mutations at codon 152 B. amyloliquefaciens subtilisin.

Figure 21 depicts the construction of single mutations at codon 156 and double mutations at codons 156 and 166 of B. amyloliquefaciens subtilisin.

Figure 22 depicts the construction of mutations at codon 217 for B. amyloliquefaciens subtilisin.

Figure 23 depicts the kcat/Km versus pH profile for mutations at codon 156 and 166 in \underline{B} . amyloliquefaciens subtilisin.

Figure 23A depicts the kcat/Km versus pH profile for mutations at codon 156 and 166 in B. amyloliquefaciens subtilisin.

Figure 24 depicts the kcat/Km versus pH profile for mutations at codon 222 in B. amyloliquefaciens subtilisin.

Figure 25 depicts the constructing mutants at codons 94, 95 and 96.

Figures 26 and 27 depict substrate specificity of various wild type and mutant subtilisins for different substrates.

Figures 28 A, B, C and D depict the effect of charge in the P-1 binding sites due to substitutions at codon 156 and 166.

Figures 29 A and B are a stereoview of the P-1 binding site of subtilisin BPN' showing a lysine P-1 substrate bound in the site in two ways. In 29A, Lysine P-1 substrate is built to form a salt bridge with a Glu at codon 156. In 29B, Lysine P-1 substrate is built to form a salt bridge with Glu at codon 166.

Figure 30 demonstrates residual enzyme activity versus temperature curves for purified wild-type (Panel A), C22/C87 (Panel B) and C24/C87 (Panel C).

Figure 31 depicts the strategy for producing point mutations in the subtilisin coding sequence by misincorporation of "-thioldeoxynucleotide triphosphates.

Figure 32 depicts the autolytic stability of purified wild type and mutant subtilisins 170E, 107V, 213R and 107V/213R at alkaline pH.

Figure 33 depicts the autolytic stability of purified wild type and mutant subtilisins V50, F50 and F50/V107/R213 at alkaline pH.

Figure 34 depicts the strategy for constructing plasmids containing random cassette mutagenesis over residues 197 through 228.

Figure 35 depicts the oligodeoxynucleotides used for random cassette mutagenesis over residues 197 through 228.

Figure 36 depicts the construction of mutants at codon 204.

Figure 37 depicts the oligodeoxynucleotides used for synthesizing mutants at codon 204.

Detailed Description

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The inventors have discovered that various single and multiple in vitro mutations involving the substitution, deletion or insertion of one or more amino acids within a non-human carbonyl hydrolase amino acid sequence can confer advantageous properties to such mutants when compared to the non-mutated carbonyl hydrolase.

Specifically, B. amyloliquefaciens subtilisin, an alkaline bacterial protease, has been mutated by modifying the DNA encoding the subtilisin to encode the substitution of one or more amino acids at various amino acid residues within the mature form of the subtilisin molecule. These in vitro mutant subtilisins have at least one property which is different when compared to the same property of the precursor subtilisin. These modified properties fall into several categories including: oxidative stability, substrate specificity, thermal stability, alkaline stability, catalytic activity, pH activity profile, resistance to proteolytic degradation, Km, kcat and Km/kcat ratio.

Carbonyl hydrolases are enzymes which hydrolyze compounds containing

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bonds in which X is oxygen or nitrogen. They include naturally-occurring carbonyl hydrolases and recombinant carbonyl hydrolases. Naturally occurring carbonyl hydrolases principally include hydrolases, e.g. lipases and peptide hydrolases, e.g. subtilisins or metalloproteases. Peptide hydrolases include α-aminoacylpeptide hydrolase, peptidylamino-acid hydrolase, acylamino hydrolase, serine carboxypeptidase, metallocarboxypeptidase, thiol proteinase, carboxylproteinase and metalloproteinase. Serine, metallo, thiol and acid proteases are included, as well as endo and exoproteases.

"Recombinant carbonyl hydrolase" refers to a carbonyl hydrolase in which the DNA sequence encoding the naturally occurring carbonyl hydrolase is modified to produce a mutant DNA sequence which encodes the substitution, insertion or deletion of one or more amino acids in the carbonyl hydrolase amino acid sequence. Suitable modification methods are disclosed herein and in EPO Publication No. 0130756 published January 9, 1985.

Subtilisins are bacterial carbonyl hydrolases which generally act to cleave peptide bonds of proteins or peptides. As used herein, "subtilisin" means a naturally occurring subtilisin or a recombinant subtilisin. A series of naturally occurring subtilisins is known to be produced and often secreted by various bacterial species. Amino acid sequences of the members of this series are not entirely homologous. However, the subtilisins in this series exhibit the same or similar type of proteolytic activity. This class of serine proteases shares a common amino acid sequence defining a catalytic triad which distinguishes them from the chymotrypsin related class of serine proteases. The subtilisins and chymotrypsin related serine proteases both have a catalytic triad comprising aspartate, histidine and serine. In the subtilisin related proteases the relative order of these amino acids, reading from the amino to carboxy terminus is aspartate-histidineserine. In the chymotrypsin related proteases the relative order, however is histidine-aspartate-serine. Thus, subtilisin herein refers to a serine protease having the catalytic triad of subtilisin related proteases.

"Recombinant subtilisin" refers to a subtilisin in which the DNA sequence encoding the subtilisin is modified to produce a mutant DNA sequence which encodes the substitution, deletion or insertion of one or more amino acids in the naturally occurring subtilisin amino acid sequence. Suitable methods to produce such modification include those disclosed herein and in EPO Publication No. 0130756. For example, the subtilisin multiple mutant herein containing the substitution of methionine at amino acid residues 50, 124 and 222 with phenylalanine, isoleucine and glutamine, respectively, can be considered to be derived from the recombinant subtilisin containing the substitution of glutamine at residue 222 (Q222) disclosed in EPO Publication No. 0130756. The multiple mutant thus is produced by the substitution of phenylalanine for methionine at residue 50 and isoleucine for methionine at residue 124 in the Q222 recombinant subtilisin.

"Carbonyl hydrolases" and their genes may be obtained from many procaryotic and eucaryotic organisms. Suitable examples of procaryotic organisms include gram negative organisms such as <u>E. coli</u> or pseudomonas and gram positive bacteria such as micrococcus or bacillus. Examples of eucaryotic organisms from which carbonyl hydrolase and their genes may be obtained include yeast such as <u>S. cerevisiae</u>, fungi such as Aspergillus sp., and non-human mammalian sources such as, for example, Bovine sp. from which the gene encoding the carbonyl hydrolase chymosin can be obtained. As with subtilisins, a series of carbonyl hydrolases can be obtained from various related species which have amino acid sequences which are not entirely homologous between the members of that series but which nevertheless exhibit the same or similar type of biological activity. Thus, non-human carbonyl hydrolase as used herein has a functional definition which refers to carbonyl hydrolases which are associated, directly or indirectly, with procaryotic and non-human eucaryotic sources.

A "carbonyl hydrolase mutant" has an amino acid sequence which is derived from the amino acid sequence of a non-human "precursor carbonyl hydrolase". The precursor carbonyl hydrolases include naturally-occurring carbonyl hydrolases and recombinant carbonyl hydrolases. The amino acid sequence of the carbonyl hydrolase mutant is "derived" from the precursor hydrolase amino acid sequence by the substitution, deletion or insertion of one or more amino acids of the precursor amino acid sequence. Such modification is of the "precursor DNA sequence" which encodes the amino acid sequence of the precursor carbonyl hydrolase rathern than manipulation of the precursor carbonyl hydrolase per se. Suitable methods for such manipulation of the precursor DNA sequence include methods disclosed herein and in EPO Publication No. 0130756.

Specific residues of <u>B. amyloliquefaciens</u> subtilisin are identified for substitution, insertion or deletion. These amino acid position numbers refer to those assigned to the <u>B. amyloliquefaciens</u> subtilisin sequence presented in Fig. 1. The invention, however, is not limited to the mutation of this particular subtilisin but extends to precursor carbonyl hydrolases containing amino acid residues which are "equivalent" to the particular identified residues in B. amyloliquefaciens subtilisin.

A residue (amino acid) of a precursor carbonyl hydrolase is equivalent to a residue of <u>B</u>. <u>amyloliquefaciens</u> subtilisin if it is either homologous (i.e., corresponding in position in either primary or tertiary structure) or analogous to a specific residue or portion of that residue in <u>B</u>. <u>amyloliquefaciens</u> subtilisin (i.e., having the same or similar functional capacity to combine, react, or interact chemically).

In order to establish homology to primary structure, the amino acid sequence of a precursor carbonyl hydrolase is directly comparted to the <u>B. amyloliquefaciens</u> subtilisin primary sequence and particularly to a set of residues known to be invariant in all subtilisins for which sequence is known (Figure 5C). After aligning the conserved residues, allowing for necessary insertions and deletions in order to maintain alignment (i.e., avoiding the elimination of conserved residues through arbitrary deletion and insertion), the residues equivalent to particular amino acids in the primary sequence of <u>B. amyloliquefaciens</u> subtilisin are defined. Alignment of conserved residues preferably should conserve 100% of such residues. However, alignment of greater than 75% or as little as 50% of conserved residues is also adequate to define equivalent residues. Conservation of the catalytic triad, Asp32/His64/Ser221 should be maintained.

For example, in Figure 5A the amino acid sequence of subtilisin from <u>B</u>. <u>amyloliquefaciens <u>B</u>. <u>subtilisin</u> var. I168 and <u>B</u>. <u>lichenformis</u> (carlsbergensis) are aligned to provide the maximum amount of homology between amino acid sequences. A comparison of these sequences shows that there are a number of conserved residues contained in each sequence. These residues are identified in Fig. 5C.</u>

These conserved residues thus may be used to define the corresponding equivalent amino acid residues of B. amyloliquefaciens subtilisin in other carbonyl hydrolases such as thermitase derived from Thermoactinomyces. These two particular sequences are aligned in Fig. 5B to produce the maximum homology of conserved residues. As can be seen there are a number of insertions and deletions in the thermitase sequence as compared to B. amyloliquefaciens subtilisin. Thus, in thermitase the equivalent amino acid of Tyr217 in B. amyloliquefaciens subtilisin is the particular lysine shown beneath Tyr217.

In Fig. 5A, the equivalent amino acid at position 217 in B. amyloliquefaciens subtilisin is Tyr. Likewise, in B. subtilis subtilisin position 217 is also occupied by Tyr but in B. licheniformis position 217 is occupied by Leu.

Thus, these particular residues in thermitase, and subtilisin from <u>B. subtilisin</u> and <u>B. licheniformis</u> may be substituted by a different amino acid to produce a mutant carbonyl hydrolase since they are equivalent in primary structure to Tyr217 in <u>B. amyloliquefaciens</u> subtilisin. Equivalent amino acids of course are not limited to those for Tyr217 but extend to any residue which is equivalent to a residue in <u>B. amyloliquefaciens</u> whether such residues are conserved or not.

Equivalent residues homologous at the level of tertiary structure for a precursor carbonyl hydrolase whose tertiary structure has been determined by x-ray crystallography, are defined as those for which the atomic coordinates of 2 or more of the main chain atoms of a particular amino acid residue of the precursor carbonyl hydrolase and B. amyloliquefaciens subtilisin (N on N, CA on CA, C on C, and O on O) are within 0.13nm and preferably 0.1nm after alignment. Alignment is achieved after the best model has been oriented and positioned to give the maximum overlap of atomic coordinates of non-hydrogen protein atoms of the carbonyl hydrolase in question to the B. amyloliquefaciens subtilisin. The best model is the crystallographic model giving the lowest R factor for experimental diffraction data at the highest resolution available.

R factor =
$$\frac{\sum |Fo(h)| - |Fc(h)|}{\sum |Fo(h)|}$$

Equivalent residues which are functionally analogous to a specific residue of <u>B. amyloliquefaciens</u> subtilisin are defined as those amino acids of the precursor carbonyl hydrolases which may adopt a conformation such that they either alter, modify or contribute to protein structure, substrate binding or catalysis in a manner defined and attributed to a specific residue of the <u>B. amyloliquefaciens</u> subtilisin as described herein. Further, they are those residues of the precursor carbonyl hydrolase (for which a tertiary structure has been obtained by x-ray crystallography), which occupy an analogous position to the extent that although the main chain atoms of the given residue may not satisfy the criteria of equivalence on the basis of occupying a homologous position, the atomic coordinates of at least two of the side chain atoms of the residue lie with 0.13nm of the corresponding side chain atoms of <u>B. amyloliquefaciens</u> subtilisin. The three dimensional structures would be aligned as outlined above.

Some of the residues identified for substitution, insertion or deletion are conserved residues whereas others are not. In the case of residues which are not conserved, the replacement of one or more amino acids is limited to substitutions which produce a mutant which has an amino acid sequence that does not correspond to one found in nature. In the case of conserved residues, such replacements should not result in a naturally occurring sequence. The carbonyl hydrolase mutants of the present invention include the mature forms of carbonyl hydrolase mutants as well as the pro- and prepro-forms of such hydrolase mutants. The prepro-forms are the preferred construction since this facilitates the expression, secretion and maturation of the carbonyl hydrolase mutants.

"Expression vector" refers to a DNA construct containing a DNA sequence which is operably linked to a suitable control sequence capable of effecting the expression of said DNA in a suitable host. Such control sequences include a promoter to effect transcription, an optional operator sequence to control such transcription, a sequence encoding suitable mRNA ribosome binding sites, and sequences which control termination of transcription and translation. The vector may be a plasmid, a phage particle, or simply a potential genomic insert. Once transformed into a suitable host, the vector may replicate and function independently of the host genome, or may, in some instances, integrate into the genome itself. In the present specification, "plasmid" and "vector" are sometimes used interchangeably as the plasmid is the most commonly used form of vector at present. However, the invention is intended to include such other forms of expression vectors which serve equivalent functions and which are, or become, known in the art.

The "host cells" used in the present invention generally are procaryotic or eucaryotic hosts which preferably have been manipulated by the methods disclosed in EPO Publication No. 0130756 to render them incapable of secreting enzymatically active endoprotease. A preferred host cell for expressing subtilisin is the Bacillus strain BG2036 which is deficient in enzymatically active neutral protease and alkaline protease (subtilisin). The construction of strain BG2036 is described in detail in EPO Publicatin No. 0130756 and further described by Yang, M.Y., et al. (1984) J. Bacteriol. 160, 15-21. Other host cells for expressing subtilisin include Bacillus subtilis 1168 (EPO Publication No. 0130756).

Host cells are transformed or transfected with vectors constructed using recombinant DNA techniques. Such transformed host cells are capable of either replicating vectors encoding the carbonyl hydrolase mutants or expressing the desired carbonyl hydrolase mutant. In the case of vectors which encode the pre or prepro form of the carbonyl hydrolase mutant, such mutants, when expressed, are typically secreted from the host cell into the host cell medium.

"Operably linked" when describing the relationship between two DNA regions simply means that they are functionally related to each other. For example, a presequence is operably linked to a peptide if it functions as a signal sequence, participating in the secretion of the mature form of the protein most probably involving cleavage of the signal sequence. A promoter is operably linked to a coding sequence if it controls the transcription of the sequence; a ribosome binding site is operably linked to a coding sequence if it is positioned so as to permit translation.

The genes encoding the naturally-occurring precursor carbonyl hydrolase may be obtained in accord with the general methods described herein in EPO publication No. 0130756.

Once the carbonyl hydrolase gene has been cloned, a number of modifications are undertaken to enhance the use of the gene beyond synthesis of the naturally-occurring precursor carbonyl hydrolase. Such modifications include the production of recombinant carbonyl hydrolases as disclosed in EPO

Publication No. 0130756 and the production of carbonyl hydrolase mutants described herein.

The carbonyl hydrolase mutants of the present invention may be generated by site specific mutagenesis (Smith, M. (1985) Ann, Rev. Genet. 423; Zoeller, M.J., et al. (1982) Nucleic Acid Res. 10, 6487-6500), cassette mutagenesis (EPO Publication No. 0130756) or random mutagenesis (Shortle, D., et al. (1985) Genetics, 110, 539; Shortle, D., et al. (1986) Proteins: Structure, Function and Genetics, 1, 81; Shortle, D. (1986) J. Cell. Biochem., 30, 281; Alber, T., et al. (1985) Proc. Natl. Acad. of Sci., 82, 747; Matsumura, M., et al. (1985) J. Biochem., 260, 15298; Liao, H., et al. (1986) Proc. Natl. Acad. of Sci., 83 576) of the cloned precursor carbonyl hydrolase. Cassette mutagenesis and the random mutagenesis method disclosed herein are preferred.

The mutant carbonyl hydrolases expressed upon transformation of suitable hosts are screened for enzymes exhibiting one or more properties which are substantially different from the properties of the precursor carbonyl hydrolases, e.g., changes in substrate specificity, oxidative stability, thermal stability, alkaline stability, resistance to proteolytic degradation, pH-activity profiles and the like.

A change in substrate specificity is defined as a difference between the kcat/Km ratio of the precursor carbonyl hydrolase and that of the hydrolase mutant. The kcat/Km ratio is a measure of catalytic efficienty. Carbonyl hydrolase mutants with increased or diminished kcat/Km ratios are described in the examples. Generally, the objective will be to secure a mutant having a greater (numerically large) kcat/Km ratio for a given substrate, thereby enabling the use of the enzyme to more efficiently act on a target substrate. A substantial change in kcat/Km ratio is preferably at least 2-fold increase or decrease. However, smaller increases or decreases in the ratio (e.g., at least 1.5-fold) are also considered substantial. An increase in kcat/Km ratio for one substrate may be accompanied by a reduction in kcat/Km ratio for another substrate. This is a shift in substrate specificity, and mutants exhibiting such shifts have utility where the precursor hydrolase is undesirable, e.g. to prevent undesired hydrolysis of a particular substrate in an admixture of substrates. Km and kcat are measured in accord with known procedures, as described in EPO Publication No. 0130756 or as described herein.

Oxidative stability is measured either by known procedures or by the methods described hereinafter. A substantial change in oxidative stability is evidenced by at least about 50% increase or decrease (preferably decrease) in the rate of loss of enzyme activity when exposed to various oxidizing conditions. Such oxidizing conditions are exposure to the organic oxidant diperdodecanoic acid (DPDA) under the conditions described in the examples.

Alkaline stability is measured either by known procedures or by the methods described herein. A substantial change in alkaline stability is evidenced by at least about a 5% or greater increase or decrease (preferably increase) in the half life of the enzymatic activity of a mutant when compared to the precursor carbonyl hydrolase. In the case of subtilisins, alkaline stability was measured as a function of autoproteolytic degradation of subtilisin at alkaline pH, e.g. for example, 0.1M sodium phosphate, pH 12 at 25 or 30 oc.

Thermal stability is measured either by known procedures or by the methods described herein. A substantial change in thermal stability is evidenced by at least about a 5% or greater increase or decrease (preferably increase) in the half-life of the catalytic activity of a mutant when exposed to a relatively high temperature and neutral pH as compared to the precursor carbonyl hydrolase. In the case of subtilisins, thermal stability is measured by the autoproteolytic degradation of subtilisin at elevated temperatures and neutral pH, e.g., for example 2mM calcium chloride, 50mM MOPS pH 7.0 at 59 °C.

The inventors have produced mutant subtilisins containing the substitution of the amino acid residues of B. amyloliquefaciens subtilisin shown in Table I. The wild type amino acid sequence and DNA sequence of B. amyloliquefaciens subtilisin is shown in Fig. 1.

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TABLE I

	Residue	Replacement Amino Acid
5	Tyr21	FA
	Thr22	С
	Ser24	С
	Asp32	QS
	Ser33	AT
10	Asp36	AG
	Gly46	V
	Ala48	EVR
	Ser49	CL .
	Met50	CFV
15	Asn77	D
	Ser87	C
	Lys94	С
	Val95	C
	Leu96	D
20	Tyr104	ACDEFGHIKLMNPQRSTVW
	lle107	V
	Gly110	CR
	Met124	IL
	Asn155	ADHQT
25	Glu156	QS
	Gly166	CEILMPSTWY
	Gly169	CDEFHIKLMNPQRTVWY
	Lys170	ER
	Tyr171	F
30	Pro172	EQ
	Phe189	ACDEGHIKLMNPQRSTVWY
1	Asp197	RA
	Met199	1
	Ser204	CRLP
35	Lys213	RT
	Tyr217	ACDEFGHIKLMNPQRSTVW
	Ser221	AC

The different amino acids substituted are represented in Table I by the following single letter designations:

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Amino acid or residue thereof	3-letter symbol	1-letter symbol
Alanine	Ala	A
Glutamate	Glu	E
Glutamine	Gln	Q
Aspartate	Asp	D
Asparagine	Asn	N
Leucine	Leu	L
Glycine	Gly	G
Lysine	Lys	` κ
Serine	Ser	S
Valine	Val	V
Arginine	Arg	R
Threonine	Thr	Т
Proline	Pro	Р
Isoleucine	lle .	1
Methionine	Met	М
Phenylalanine	Phe	F
Tyrosine	Tyr	Υ
Cysteine	Cys	С
Tryptophan	Trp	W
Histidine	His	Н

Except where otherwise indicated by context, wild-type amino acids are represented by the above three-letter symbols and replaced amino acids by the above single-letter symbols. Thus, if the methionine at residue 50 in B. amyloliquefaciens subtilisin is replaced by phenylalanine, this mutation (mutant) may be designated Met50F or F50. Similar designations are used for multiple mutants.

In addition to the amino acids used to replace the residues disclosed in Table I, other replacements of amino acids at these residues are expected to produce mutant subtilisins having useful properties. These residues and replacement amino acids are shown in Table II.

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TABLE II

Residue	Replacement Amino Acid(s)
Tyr-21	L
Thr22	K
Ser24	A
Asp32	
Ser33	G
Gly46	
Ala48	
Ser49	
Met50	LKIV
Asn77	D
Ser87	N
Lys94	RQ
Val95	LI
Tyr104	
Met124	KA
Ala152	CLITM
Asn155	
Glu156	ATMLY
Gly166	
Gly169	
Tyr171	KREQ
Pro172	DN
Phe189	
Tyr217	
Ser221	
Met222	

Each of the mutant subtilisins in Table I contain the replacement of a single residue of the <u>B</u>. <u>amyloliquefaciens</u> amino acid sequence. These particular residues were chosen to probe the influence of such substitutions on various properties of B. amyloliquefacien subtilisin.

Thus, the inventors have identified Met124 and Met222 as important residues which if substituted with another amino acid produce a mutant subtilisin with enhanced oxidative stability. For Met124, Leu and Ile are preferred replacement amino acids. Preferred amino acids for replacement of Met222 are disclosed in EPO Publication No. 0130756.

Various other specific residues have also been identified as being important with regard to substrate specificity. These residues include Tyr104, Ala152, Glu156, Gly166, Gly169, Phe189 and Tyr217 for which mutants containing the various replacement amino acids presented in Table I have already been made, as well as other residues presented below for which mutants have yet to be made.

The identification of these residues, including those yet to be mutated, is based on the inventors' high resolution crystal structure of B. amyloliquefaciens subtilisin to 1.8 A (see Table III), their experience with in vitro mutagenesis of subtilisin and the literature on subtilisin. This work and the x-ray crystal structures of subtilisin containing covalently bound peptide inhibitors (Robertus, J.D., et al. (1972) Biochemistry 11, 2439-2449), product complexes (Robertus, J.D., et al. (1972) Biochemistry 11, 4293-4303), and transition state analogs (Matthews, D.A., et al. (1975) J. Biol. Chem. 250, 7120-7126; Poulos, T.L., et al. (1976) J. Biol. Chem. 251, 1097-1103), has helped in identifying an extended peptide binding cleft in subtilisin. This substrate binding cleft together with substrate is schematically diagramemed in Fig. 2, according to the nomenclature of Schechter, I., et al. (1967) Biochem Bio. Res. Commun. 27, 157. The scissile bond in the substrate is identified by an arrow. The P and P' designations refer to the amino acids which are positioned respectively toward the amino or carboxy terminus relative to the scissle bond. The S and S' designations refer to subsites in the substrate binding cleft of subtilisin which interact with the corresponding substrate amino acid residues.

Atomic Coordinates for the Apoenzyme Form of B, Amyloliquefaciens Subtilisin to 1.8AResolution

5										
			19.434	\$3.195	-21.756	3	ALE CE	29.811		- 34 - 44
	1	ALA C	14.731	\$0.125	-21.324	í	414 0	10.374	\$1.174 \$1.197	-21.965 -28.175
	;	ALA CB	23.099	51.518	-21.183	i	51.4 4	10.265	49.884	-22.941
	į	GL & CA	17.217	49.000	-21.434	;	GLEC	17.875	47.704	
	2	610 0	10.745	47.145	-21.691	2	SLE CO	14.125		-20.992
	•	610 66	15.324	47.905	-21.927	2	STH CD	13.912	48.740	-22.449
10	ź	CL = DE1	13.021	48.412	-22.847	ź	era mes	14.115	47.742	-22-930
70	5			47.205		i	SER CA		44-917	-23.926
	_	TER C	17.477	44.928	+19.852 -19.440	;	SEP O	17.930 15.390	45.141	-19.437
	3	358 68	16.540	45.038	-18.047	;	St# 06	17.482	45.352	-19.229
	•	VAL H	14.991	43.444	-19.725		VAL CA	15.944	46.218	-17.049 -19.439
	- 1	VAL E	14.127	43.934	-18.290		VAL D	17.123	41-170	
	- 7	VAL CO	14.008	41.622	-20.022	i	VAL CGI	14.874	48.572	-18.006 -28.741
	- ;	VAL CGZ	14.037	42.246	-22.106	š	PRO te	15.239	42.104	-17.331
15	;	PPD CA	15.384	41.415	-14.027	ś	PRD C	15.301	39.905	-16.249
	í	PRO O	14.885	39.243	-17.146	5	PRO CS	14.150	41.880	-15.26)
	į	P80 EG	13.241	43.215	-15.921	5	PIO CO	24.944	42.786	-17.417
	í	TTR N	14.343	37.240	-15.687	í	TTR CA	16.624	37.803	-15.715
	- ĕ	TYR C	15.359	34.975	-15.528	ě	TTE D	15.224	35.743	-14.235
	- 4	118 68	17.824	37.323	-14.834	ĭ	TTR CG	18.021	35.047	-15.055
	ě	TTR CD1	18.437	35.452	-16.346	ě	TYR CD2	17.496	34.904	-14.071
	ě	TYR CEL	10.535	34.070	-14.453	ě	TTE CEZ	17.015	33.539	-14.379
20	ă	TTR CI	18.222	33.154	-15.628	6	TTR OH	10.312	31.830	-15.996
	7	GLT B	14.464	37.362	-14-630	7	GLY CA	13.211	34.440	-14.376
	7	GLT C	12.400	34.535	-15.670	7	GLT D	11.747	35.478	-15.883
	•	VAL M	12.441	37.329	-14.541	4	VAL CA	21.777	37.523	-17.636
		VAL C	12.343	34.433	-10.735		VAL O	11.639	35.716	-11.470
		TAL CA	11.765	34.900	-10.567		VAL CG1	11.104	31.693	-11.943
	ı	TAL CG2	18.991	39.419	-17.733	•	SER M	13.661	36.310	-10.775
25	9	SER CA	14.419	35.342	-17.562	9	SEA C	14.100	33.720	-18.945
	9	\$ E B O	14.112	33.014	-19.301	•	SEA CO	85.924	35.432	-19.505
		SEE DC	16.162	34.747	-20.358	10	CLM #	84.115	33.017	-17.462
	10	GLN CA	13.964	32.636	-16.876	10	STM C	12.607	31.417	-17.277
	10	GLW D	12.745	30.442	-17.413	2.0	ern CB	14.125	32.115	-15.410
	30	ELM CC	14.295	31.617	-14.580	10	GTM CD	14.484	31.911	-13.147
	10	ern DEI	14.554	33.040	-12-744	10	PIN NES	14.552	30-940	-12.251
	11	ILE N	11.625	32.575	-17.470	31	ILE CA	10.373	31.904	-18.102
30	11	ILE C	10.209	31.792	-19.405	11	Itt O	9.173	31.353	-20.100
	11	116 CES	9.132	32.469	-17.475	11	ILE CG1	1.044	34.117	-16.049
	12	LTS R	9.142 11.272	32.455	-15.941 -20.277	11 12	ILE COI	7.588	34.640	-17.923
	12	145 6	30.454	33.004	-22.522	12	L75 D	11.300 10.178	32.111	-21.722 -23.606
	12	LTS CO	31.257	30.444	-22.216	12	LYS CG	12.283	32.783	
	ii	LYS CO	12.543	28.517	-22.159	12	LYS CE	13.023	27.467	-21.423 -21.166
	12	LYS MI	14.474	27.680	-20.935	ii	ALA R	10.109	34.130	-21.991
35	13	ALA CA	9.325	35-198	-22.431	11	ALA C	10.024	35.716	-23.863
	11	ALA D	9.336	35.804	-24.901	13	ALA CO	0.005	16.193	-21.565
	14	PED W	11.332	35.150	-23.893	14	PRO CA	11.985	34.430	-25.120
	1.	>40 C	11.786	35.357	-24.317	14	P20 D	11.778	36.047	-27.445
	14	P#0 C#	13.462	34.510	-24.692	14	PED CC	13.324	36.978	-23.221
	14	PED CD	32.241	35.736	-22.758	15	ALA M	11.540	34.236	-26.129
	15	ALA CA	21.379	33.450	-27.367	15	ALA C	10.002	33.795	-28.032
40	15	ALA O	10.001	33.710	-29.278	15	ALG CS	11.552	31.969	-21.042
40	16	LEU m	9.085	34.138	-27.240	16	LEU CA	7.791	34.558	-27.020
	14	LEU C	7.912	31.925	-24.521	14	rin o	7.342	34.114	-29.568
	16	LEU CB	6.746	34.423	-24.698	16	ren ce	3.790	33.465	-24.522
	14	L EU COL	3.881	33.234	-27.009	14	FED CDS	4.694	32.207	-24.283
	17	#15 b	8.445	34.070	-27.922	37	MIS CA	0.890	30.151	-20.531
	17	MIS C	9.510	37.101	-29.898	17	HIS D	9.107	30.672	-10.854
	17	W15 CB W15 @81	9.701	39.106	-27.652	17 17	MIS CC	9.195	19.201	-24.242
45	17	MIS CES	9.938 9.224	39.887	-25.272	17	MIS COZ		38.924	-Z5.694
	19	. 324 0	10.44)	37-833	-24.144 -38.022	10	SEE CA	8.079 11.109	39.328 34.739	-34.381 -31.322
						• •	•=			

EE

		BER C	40						
	11		10-117	86-123	-32.353	34 SE* D	30.947	\$6.132	-33.034
	14	168 68	11.111	35.711	-31.372	38 500 95	13.371	36.450	-34.311
	19	ALM M	9.000	\$5.495	-31.843	37 6L% C4	0.912	84.942	-32.871
	19	GLM E	7.142	36.111	-33.303	19 6LM D	4.211	\$1.972	-34-219
	19	BLW CS	7.221	33.741	-32.280	19 BL4 [6	7.913	22.602	-31.621
_	11	8L# CD	6.923	\$1.701	-31.181	- 37 6L% BE1	5.719	31.433	-31.444
5	3.0	SLM MEZ	7.342	\$0.052	-30.234	80 SLT N	7.200	37.223	-32.887
	10	BLT CA	4.349	30.307	-32.159	20 617 C	5.101	30.492	-31.000
	20	SLT D	4.243	31.274	-32.215	21 178 6	\$.202		
	ii	778 C4	4.116		-21.763			37.801	-38.741
	ii	TYR D		37.431		83 TYR C	4.879	31.552	-20.525
	11	778 CG	8.422	31.074	-27.766	83 TT4 CE	3.471	34.471	-29.443
			2.973	31.784	-30.100	21 TT# CO1	1.795	34.331	-31.230
	33	144 CDS	3.450	34.794	-11.597	21 TYP CE1	1.306	\$3.797	-32.445
10	31	148 645	1.173	34.241	-32.000	21 440 61	2.003	34.755	-33.047
_	21	TYR OR	1.503	34.241	-34.250	22 THE N	3.992	37.680	-28.284
	22	THE CA	4.242	40.327	-27.129	SS THE C	3.071	41.922	-24.244
	11	7#8 G	3.287	41.725	-25.325	22 445 68	9.173	41.751	-27.411
	22	THE DG1	4.319	42.457	-28.597	23 PMP C62	6.474	41.323	-20.219
	21	SLT M	1.737	40.285	-24.453	23 6L7 CA	0.000	49.400	-23.542
	23	SLT E	-0.197	41.431	-26.118	23 BLT D	-1.013	42.075	-29.310
	24	114 W	-0.023	41.947	-27.871	24 550 60	-0.017	41.917	-20.012
15	24	SER C	-2.141	41.626	-27.864	24 119 0	-2.612	41.000	
75	24	SER CO	-0.734	41.125	-29.520	24 588 06	0.163	43.432	-20.100
	25	45+ 4	-3.059	47.412	-27.510	28 ASH CA	-4.519	43.487	-20.728
	11	414 6	-9.018	42.073	-24.203	25 ASN D	-6.233		-27.313
	ží	ASA CB	-0.165	43.227	-20.700	21 A3h C4	-4.940	42.661	-24-119
	11	ASE ODI	-4.745	43.747	-31.003			44-170	-29.005
	ii	TAL M	-4.177	41.449	-23.292	25 ASH 002 26 VAL CA	-4.747	45.461	-29.854
	11	VAL C	-4.792	42.452	-22.007		-4,474	41.479	-24.143
	11	VAL ES	-3.714	40.503	-23.021	86 VAL D	-1.010	43.419	-21.611
20	1	VAL CEZ	-3.111			24 VAL C61	-4.140	39.402	-22.548
	27	LTS CA	-6.113	34.574	-20.018	81 142 4	-3.910	42.613	-22.301
	1,	LY 1 0		43.524	-21.175	27 LTE C	-1.915	42.872	-19.041
	27	111 66	-4.405	41.073	-17.413	27 LTS C8	-7.990	43.912	-21.249
	27	LTS CE	-8.044	44.573	-22.490	33 FAR CD	-1.321	49.301	-21.610
			-10.304	49.497	-23.137	\$7 LYS #2	-9.616	44.251	-24.244
		VAL N	-4.318	43.442	-19.200	SO ANT CO	-4.487	42.988	-27.897
	31	447 6	-4.750	43.131	-14.828	SE ANT D	-4.204	45.095	-14.817
25	31	TAL CO	-2.124	42.444	-17.932	30 AAF C21	-2.404	42.193	-14.889
	11	ANT CES	-2.667	41.805	-19.173	II ALE W	-3.414	43.527	-19.813
	31	ALA CA	-8.747	44.330	-14.639	29 ALB C	-4.750	44.010	-13.91)
	31	ALA D	-4.446	42.845	-13.104	89 ALA CB	-7.172	44.187	-14.181
	30	VAL B	-4.057	45.033	-13.072	BO AFF CV	-1.144	44.942	-11.910
	30	VAL C	-3.956	45.409	-10.681	BD VAL D	-4.155	44.648	-10.876
	30	WAL CB	-1.884	49.810	-12.149	30 VAL CG1	-0.9,54	45.901	-18.980
	30	ANT COS	-2.85)	48.234	-13.357	31 ILF W	-4.914	44.518	-9.877
30	31	ILE CA	-0.324	44.846	-8.679	31 ILE C	-4.344	44.113	-7.548
	31	ILE O	-1.825	43.915	-6.997	31 314 CF	-6.457	43.774	-8.901
	31	368 C61	-7.278	43.707	-9.791	D1 IL8 C62	-7.274	44.836	-7.221
	31	ILT COI	-8.617	41.954	-9.717	32 457 4	-4.044	44.113	-7.217
	75	43 P CA	-2.944	44.467	-4.255	32 489 C	-3.071	47.889	-3.785
	35	ALP D	-4.197	48.418	-8.362	32 48º E6	-1.475	44.129	-1.092
	31	41° C6	-1.413	48.782	-6.273	31 43° 001	8.934	44.592	-4.876
	3 2	48 903	-6.881	44.429	-1.330	33 844 0	-1.731	40.012	-1.314
35	31	51 F C4	-1.895	48.857	-4.801	31 11° C	-3.952	90.174	-3.888
	33	SER D	-1.704	\$2.134	-1.343	33 \$11 61	-0.621	49.922	-3.939
	22	30 4 06	8.331	80.026	-4.774	34 BLT W	-2.173	81.740	-1.11
	34	GLT CA	-1.111	81.728	-9.141	34 8L7 E	-1.635	81.648	-9.817
	34	SLT D	-8.144	80.031	-0.761	DF ELE A	-0.765	82.471	-10.102
	31	ILE CA	0.200	\$2.414	-10.975	Di ili i	0.566	83.919	-11.243
	33	ILE D	-0.327	14.431	-11.744	Di ili co	-0.0.2	D1.094	-12.367
	11	JLF CGS	-0.530	60.210	-12.091	30 \$1.2 662	1.149	\$1.741	-13.307
40	33	ILF COL	-0.942	47.411	-13.424	31 437 1	1.616	14.253	-10.971
	14	ASP CA	2.311	85.618	-11.232	34 450 5	2.201	31.014	-12.702

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	3 6	45 0	3.004	\$5.471	-13.579	34	ASP CB	3.712	55.720	-14.514
	31	41P C6	4.339	\$7.099	-10.004	34	ASP 001	3.755	\$7.974	-11.429
		ASP 802	3.441	\$7.217	-10.243	37	510 B	3.364	54.622	-13.111
	34	168 CA	1.103	\$7.221	-14.512	37	Ste C	2.317	30.073	-14,949
	37	•	2.545	58.301	-14.151	17	Ste Co	-0.893	58.847	-14.784
	37	\$ E # D		59.113	-13.079	10	51# W	3.143	58.614	-14.00)
5	37	SER DE	-0.030		-14.487	51	SER C	5.444	\$8.705	-14.992
	31	SER CA	4.261	\$9.305		38	51 F C0	4.742	40.433	
	30	SER D	4.543	\$9.251	-15.205	31	M15 M	5.454	\$7.398	-13.341
	31	SER DG	5.374	59.865	-12.234	31	MIS C	4.401		-14.892
	37	MIS CA	4.437	\$6.574	-15.791				\$6.401	-14.774
	39	wis D	5.730	85.078	-17.419	37	M12 CB	6-437	\$5.203	-14.515
	37	M13 CE	8.414	54.607	-14.456	31	MIS MD3	0.795	54.354	-15.561
	31	MIS CD2	8.747	\$4.345	-13.369	39	M15 CE1	9.970	53.930	-15.130
10	3+	MIS MEZ	7.986	53.910	-13.608	40	P P D	7.007	54.834	-17.367
70	4.	PED CA	7.913	\$6.697	-18.831	40	PRO C	0.154	55.280	-14.357
	4.	PRO D	0.032	\$5.897	-20.374	40	PED CS	9.247	\$7.533	-19.161
	4.	PEO CG	10.053	57.485	-17-902	40	P40 CD	0.711	\$7.452	-16.774
	4.2	ASP W	0.461	54.328	-18.485	41	ASP DD2	11.140	58.399	-10.448
	41	ASP BD1	10.325	31.395	-20.429	41	ASP CC	10.473	51.307	-19.211
	4 3	ASP CO	9.777	52.239	-18.224	41	ASP CA	8.445	\$2.959	-10.764
	41	ASP C	7.311	\$2.143	-14-639	41	ASP D	7.394	50.947	-18.977
15	42	LEU M	6.185	\$2.803	-10.558	42	LEU CA	4.892	82-147	-18.466
,,,	4.2	LEU C	3.974	\$2.907	-19.374	42	LEU D	3.77)	54.163	-19.499
	4.2	LEU CB	4.471	\$2,150	-17.000	42	LEU CG	5.182	\$1.343	-15.944
	42	LEU CD1	4.535	52.546	-14.591	42	LEU CD2	5.273	41.877	-14.358
	43	LYS M	3.618	\$2.135	-19.946	43	LTS CA	1.013	52.485	-20.721
	43	LYS C	0.437	32.156	-20.010	43	L12 D	0.504	\$8.920	-19.620
	43	LTS CB	2.071	52.319	-22.169	43	LIS CE	0.615	\$2.436	-22.910
	4)	LYS CO	0.111	\$2.842	-24.339	43	LTS CE	-9.100	52.544	-25.260
20	43	LVS WZ	0.337	\$1.757	-24.418	44	VAL M	-0.171	53.835	-19.490
	44	VAL CA	-1.407	\$2.437	-19.765	44	ANT C	-2.571	52.887	-19.731
	44	TAL D	-2.623	53,786	-28.434	44	VAL EB	-1.480	53.351	-17.343
	4.4	VAL CGI	-2.724	52.941	-14.502	44	VAL EEZ	-0.197	53.194	-14.553
	4.5	ALA M	-3.444	\$1.951	-19.871	45	ALA CA	-4.619	51.177	-20.810
	45	ALA C	-5.841	52.507	-20.053	45	ALA O	-4.783	\$3.015	-20.763
	4.5	ALA CB	-4.031	\$0.580	-21.309	44	SLT M	-5.910	\$2.354	-18.748
	4.6	GLY CA	-7.012	\$2.037	-11.001	46	SLY C	-6.987	\$2.443	-14.538
25	44	GLT D	-5.934	32.804	-14.035	47	SLT M	-8.892	32.438	-15.793
	47	GLT CA	-8.014	\$2.244	-14.388	47	SLT C	-9.179	52.757	-13.572
	47	SLY O	-9.944	\$1.483	-14.185	48	ALA W	-9.221	\$2.444	-12.330
	41	ALA CA	-10.235	52.070	-11.302	48	ALA C	-9.790	\$2.475	-9.741
	41	ALA D	-1.044	52.720	-9.725	48	ALA CE	-21.550	\$2.100	-11.617
	49	5 E R .	-18-149	\$3.547	-9.037	49	SER CA	-9.752	53.355	-7.452
	47	SER C	-10.947	52.986	-4.783	43	SER B	-11.972	\$3.477	-4.904
	49	SER CA	-9.092	34.500	-7.029	49	SEE BC	-8.277	\$4.255	-5.450
30	50	MET B	-10.835	\$2.007	-5.932	50	MET CA	-11.052	\$1.549	-4.974
	50	MET C	-11.463	\$1.942	-3.541	50	MET O	-11.997	51.390	-2.575
	50	MET CO	-12.017	\$0.015	-4.994	38	MET CG	-11.912	41-453	-6.317
	5.0	MET SO	-11.440	47.887	-7.256	5.0	MET CE	-12.808	50.111	-8.703
	51	TAL B	-10.477	52.744	-3.422	51	TAL CA	-7.161	\$3.170	-2.967
	51	VAL E	-18.430	\$4.542	-1.907	51	VAL D	-10.237	\$5.437	-2.402
	\$1	TAL CO	-8.443	\$3.155	-2.000	51	VAL CEL	-7.892	\$3.579	-4.631
	51	TAL CG2	-7.744	\$1.615	-2.302	52	PRD 8	-11.421	\$4.473	-1.056
35	52	PRO CA	-12.372	\$5.933	-0.821	52	POD C	-11.490	\$7.12)	-1.441
	52	PRO D	-11.771	30.220	-0.925	\$2	PRO Ca	-13.400	35.574	0.244
	52	PED CE	-13.513	\$4.183	0.015	52	PRD CO	-12.244	\$3.629	-0.175
	13	Sta m	-18.442	\$6.904	0.291	53	SER CA	-9.530	\$7.982	0.482
	11	SER C	-0.420	\$8.245	-0.324	53	548 G	-7.479	\$9.224	-4.438
	1)	\$68 CB	-1.114	87.707	2.849	\$3	518 OC	-6.254	\$4.521	2.127
	94	SLU H	-8.254	\$7.523	-1.393	54	ELU CA	-7.104	87.648	-2.421
40	54	GLU C	-7.767	\$7.303	-3.785	34	GLU 0	-7.511	86.243	-4.379
40	54	GLU CB	-6.134	56.571	-2.154	54	GLU CC	-1.211	36.959	-8.927
	44	GL# CH	-4.044	44.847	-9.478	44	CIH AFT	-1.44	81.694	-1.946

	54	ELV SEZ	-3.900	55.777	0.271	55	THE M	-0.571	98.291	-4.249
	5.5	THE CA	-9.433	58.121	-5.441	55	THE C	-8.744	50.119	-4.779
	55	THE B	-7.433	\$7.919	-7.816	\$ 5	THR CO	-10.986	\$9.280	
	35	THE OCL	-9.885	40.510	-5.418	83	THR C62	-11-432	\$9.143	-3.303
	34	ASH H	-7.482	\$8.403	-4.877	54	410 002	-4.939	41.179	-4.817
_	54	A58 001	-5.075	31.967	-10.337	54	ASH CC	-5.273		-9.881
5	54	ASS CO	-3.878	31.474	-0.208	54	ASO CA		\$9.425	-9.555
	34	459 6	-4.812	\$7.094	-8.305	56	ASH D	-6.762	\$8.425	-1.200
	57	PRO H	-4.342	\$4.261	-9.250	57		-5.184	54.944	-7.478
	37	#20 CD	-7.384	54.433	-18.272		PRD C6	-7-123	35.257	-11.177
		PRO CA	-5.679			37	PRO CO	-4-644	54.178	~10.235
	\$7			\$4.941	-9.332	57	\$10 C	-4.301	55.062	-9.944
	37	PRO D	~3.509	\$4.120	-9.945	5.0	PHE M	-3.998	36.262	-10.491
10	51	PHE CA	-2.747	54.577	-11.222	51	PREC	-1.712	37.129	-10.253
10	5.8	PHE O	-0.635	\$7.497	-10.600	54	PHE CS	-2.943	\$7.502	-12.423
	54	PHE CG	-3.983	54.748	-13.357	5.0	PHE COL	-3.756	\$5.78t	-14.859
	54	BHE COS	-5.211	57-630	-13.459	54	PHE CEL	-4-722	\$5.255	-14.924
	5.0	PHE CE2	-6.394	57.095	-14.274	51	PHE CZ	-5.947	\$5.939	-15.051
	59	STR B	-2.044	57.119	-8.998	57	GL# CA	-1.172	\$7.583	-7.934
	59	SLM C	-8.807	\$6.403	-7.800	59	GLW D	-1.439	\$4.083	-4.115
	59	GLN CB	-1.462	58.668	-7.889	59	ELW CG	-0.942	\$9.261	-4.034
15	39	GLM CD	-1.750	60.157	-5.150	51	SLE DEI	-1.404	61.288	-4.836
7.5	59	CID DES	-2.959	59.685	-4.742	60	ASP ME	0.410	\$5.095	-7.211
	40	ASP CA	0.851	\$4.792	-6.304	4.0	ASP E	1.431	\$5.267	-5.090
	40	ASP O	. 2.827	\$5.550	-5.231	63	ASP CB	1.576	\$3.744	-7.108
	40	ASP EG	2.077	\$2.534	-4.300	4.0	ASP BD1	1.744	\$2.337	-5.190
	40	85P 002	2.915	\$1.041	-7.030	41	ASM M	0.757	55.245	-3.750
	61	ASU MOZ	-1.364	\$7.747	-2.347	41	ASH DOL	0.666	\$8.544	-2.875
	41	ASH CC	-0.048	57.670	-2.399	61	ASH CB	0.531	54.463	-1-704
20	61	ASH CA	1.557	\$5.734	-2.700	61	ASH C	2.291	\$4.632	-1.940
	41	ASH D	2.733	\$4.862	-8.902	62	ASH U	2.210	53.434	-2.449
	42	ASH CA	2.877	52.348	-1.709	62	ASH C	4.124	\$1.013	-2.479
	42	ASR D	4.951	\$1.313	-1.770	62	ASW CB	1.703	\$1.319	
	4.2	ASR CG	2.371	50.103	-0.697	42	ASH OD1	2.633	49.077	-2.421
	62	45 B BO2	2.422	50.204	0.601	43	SER W	4.152		-1.343
	63	SED CO	5.169	\$1.494	-4.709	43	SER C		52.164	-3.741
	63	SEE D	5.513	49.790	-4.249	63	380 CB	5.671	54.256	-3.269
25	63	SER 06	4.871	50.494	-3.416	- 11	MIS M	4.523 4.202	\$1.950	-4.012
	44	MES CA	3. 994	48.855	-4.935		213 C		49.475	-4.439
	64	M15 0	3.061	44.974	-7.104		MIS CO	3.364	47.759	-6.261
	64	MIS CG	3.144	44.921	-3.724		WIS MD1	3.104	47.501	-3.747
	64	MIS COZ	4.054	45.194	-3.135		MIS CEI	2.107	45.247	-4.241
	44	MIS WEZ	3.556	43.920	-3.361	43	CLA M	2.414	43.944	-4.054
	45	SLT CA	1.552	40.264	-7.030	45	SLT C	2-207	48.428	-6.587
	45	617 0	2.230	48.678	-10.134			2-372	48.636	-9.037
30	64	THE CA	4.044	50.117	-9.954	44	THE M	3.233	49.659	-8.832
	66	THE D	5.333	48.789		44	THE C	5.019.	49.809	-10.291
		148 BG1	3.637		-11-461	44	THP CS	4.744	51.511	-9.667
	67	MIS D	3.405	\$2.425	-7-406	46	THR CG2	5.534	52.478	-10.047
	47	WIS C	6.091	40.443	-9.274	67	MIS CA	4.793	47.341	-7.458
	47	MIS CO	7.308	46.141	-10.143	47	M15 0	6.647	45.438	-11.150
	67				-1.044	•7	HIS CC	6.575	44.275	-0.148
35		MIS BOL	8.590	44.987	-8.276	47	MI2 CDS	9.964	46.478	-8.076
00	67	WIS CEL	9.857	44.491	-0.299	67	HIS WEL	10.478	45.514	-0.104
			4.092	45.749	-9.731	4.0	VAL CA	4.142	44.687	-10.266
	44	VAL C	3.054	44.940	-11.740	40	VAL D	4.114	43.942	-12.535
	61		2.939	44.252	-7.386	6.8	ANT CC3	1. ***	43-240	-10.020
		447 CES	3.319	43.705	-0.000	47	ALA U	3.373	46.947	-12.113
	69	ALA EA	3.637	44.444	-13.429	49	ALA C	4.313	44.370	-14.411
	67	ALA D	4.020	45.911	-15.565	44	ALA CS	2.332	67.853	-13.386
40	70	SLT R	5.340	44.707	-13.914	70	SLT CO	4.175	46.005	-14.870
	-	SLT C	7.144	45.370	-15.021	7.0	CLT D	T. 684	43.154	-14.119
	71	Tet m	4.870	44.431	-14.134	71	THE CA	7.177	43.019	-14.444
	71 71	Tak C	4-224	42.504	-15.54)	73	THE D	4.602	41.828	-16.475
	7 I	TAR CO	7.119	42.870	-13.141	71	THE BC1	0.191	42.592	-12.390

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	71	THE CES	7.274	40.583	-13.594	72 VAL B	4.936	42.887	-15.427
	72	VAL CA	3.974	42.473	-16.484	72 VAL C	4.312	43.084	-17.831
	72	VAL B	4.341	42.340	-14.840	72 VAL CO	2.914	42.847	-14.005
	72	VAL CEI	1.512	42.499	-17.170	TE VAL CEZ	2.142	42.327	
			6.50	44.417		TO BLA CA			-14.723
5	73	ALA W			-17-140		4-987	43.001	-19.167
	73	ALA C	5.433	46.333	-19.355	73 ALB D	3.042	47.188	-20.216
	73	ALA EB	3.107	45.443	-19.433	T4 ALS W	4.344	44.429	-10.635
	74	ALA CA	7.478	67.591	~18.959	76 ALR C	7.740	47.649	-20.342
	74	ALA 8	1.750	46.640	-21.054	76 ALA CO	0.433	47.444	-17.125
	75	LEU W	7.650	48.784	-21.839	75 LEU CA	7.012	41.761	-22.454
	75	LEU C	9.192	48.568	-22.966	75 LEU 0	10.142	44.750	
	75	LEU CO	7.548	\$0.471	-22.809	75 LED CG	6.123		-27.253
		LEU CDI	6.079			75 160 (82		50.913	-21.378
10	75			\$2.434	-22.300		5.014	30.462	-23.485
	74	ASH W	9.147	48.103	-24.169	76 ESM 802	12.385	46.432	-24.364
	74	ASM DDI	10.950	45.940	-27.928	74 ASH CG	11.195	44.274	-26.862
	74	ASH CB	30.010	46-651	-25.908	76 , 45M CA	18.359	47.738	-24.938
	76	ASH C	10.783	41.941	-25.643	76 ASH O	30.157	49.479	-26.619
	77	ASE B	11.004	41.444	-23.071	TT ASH CA	12.220	\$0.957	-25.681
	77	ASH C	13.707	51.029	-23.348	TT ASW D	14.364	49.979	-25.319
	77	ASU CO	11.335	\$2.076	-25.117	TT ASM CG	11.250	\$2.027	-23.616
46	77	434 OD1	12.032	\$1.344	-22.917	TT ASH ND2	10.294	52.741	-23.025
15	78	SER R	14.125	\$2.267	-25.144	78 SER CA	15.513		
								\$2.614	-24.906
	78		15.818	\$2.742	-23.434		14.902	53.071	-23.164
	70	SER CB	15.905	53.941	-25.517	78 SER D6	15.926	53.870	-26.999
	79	ILE M	14.858	52.565	-22.529	TO ILE CA	15.155	\$2.784	-21.120
	79	ILE C	14.617	51.483	-20.230	79 ILE D	13.843	30.041	-28.679
	79	STE CO	14.471	54_174	-20-017	79 ILE C61	12.945	54.032	-28.814
	79	ILE CG2	34.997	\$5.320	-21.612	79 ILE CO1	12.135	\$5.174	-20.155
20		GLT M	14.995	51.768	-10.941	BO ELT CA	14.476	50.949	-17.913
	80	SLT C	14.412	47.448	-18.219	SO ELY D	15.719	48.994	-18.544
	01	VAL W	13.513	48.766	-17.980	81 VAL CA	13.411	47.284	-18.041
	81	VAL C	12.511	44.717	-19.217	81 VAL D	12.260	47.739	-20-117
	i	VAL CO	13.001	44.755	-14-677	BI VAL CEI	14.930	47.004	-15-573
	. i	VAL CG2	11-638	47.261	-16.231	82 LEU N	12.126		
		LEU CA		45.020				45.445	-19.216
	8 Z		11.312		-20.256	B2 LEU C	10.376	44.028	-19.510
25	8.5	LEU O	10.451	43.356	-18.600	#2 LEU CB	12.204	44.219	-21.229
25	82	FER CE	11.430	43.561	-22.366	MZ LEU COL	28.796	44.657	-23-223
	15	TEO CDS	12.359	42.675	-23.192	13 GLY W	9-131	44.180	-19.816
	0.3	GLT CA	4.133	43.323	-19.114	83 GLT C	6.627	42.011	-14.525
	93	GLT G	8.544	41.822	-21.026	Se VAL N	7.272	41.112	-14.243
	84	ANT CV	4.973	39.807	-19.888	B4 VAL C	4.164	48.830	-21.140
	84	TAL D	4.424	39.472	-22.194	D4 VAL CB	6.256	36.920	-14.441
		TAL CGS	5.400	37.677	-19.557	64 WAL EG2	7.190	38.507	-17.705
	85	ALA M	5.154	40.924	-21.024	US ALA CA	4.217	41.194	-22.158
30	95	ALB C	4.213	42-483	-22.396	#5 ALA D	3.240	43.401	-22.030
	85	ALA CO	2.846	40.443	-21.748	86 PPD #	5.240	43.180	
	86	PED CA	5.413	44.435	-23.205	84 P80 C	4.321		-23.859
	0.4	P40 0	4.291	46.605	-23.849	86 PRD CB		45.371	-23.947
	16	P#0 C6					6.522	44.784	-23.413
			7.030	43.446	-24.546	84 PED CD	6.377	42.440	~23.436
	47	SER W	3.548	44-676	-24.769	ST SER CA	2.489	45.324	-25.529
	87	24 C	1.103	45.332	-24.871	87 SEP 0	0.142	45.513	-25.619
35	87	SER CO	2.401	44.377	-24.921	47 SER 05	3.591	45.143	-27.583
	**	ALA M	1.017	44,584	-23.742	88 ALA CT	-0.143	43.510	-21.628
	••	ALA CA	-8.273	44.353	-23.084	PD ALA C	-0.811	45.717	-22.690
		44 8	-0.374	46.717	-22.435	to See n	-2-219	41.691	-22.678
		SER 05	-4.144	47.302	-24.280	89 SER CS	-4.343	44,793	-22.016
	87	SER CA	-3.801	44.067	-22.221	89 SER C	-3-134	46.780	-20.727
	-	3 E P O	-3.193	45.844	-20.209	10 Ltu m	-2-446	47.656	-20.037
	98	LEU CA	-2.178	47.667	-18.593	10 LEU C	-1.483	48.438	
40	90	Ltu B	-3.502	49.604	-18.215	10 LEU CS			-17.864
₩0	**	Ltu C6	-0.233	47.831	-17.17-		-0.731	48.273	-18.426
	70	14n C03					-0.626	44.341	-17-219
			1.160	40.124	-17.047	91 TYR W	-4.264	47.144	-14.938
	91	TTT CA	-3.258	48.678	-16.137	41 AAB C	-4.873	40.730	-14.485

	91	***	-4.474	47.749	-14.073	91	TTO CO	-4.684	48.093	- 6 4 9 4 4 4
	91	TYR C4	-1.494	48.237	-17.741	93	TTR CD1			-14.314
			-7.971					-4.595	47.415	-10.755
	91	TER COS		49.275	-10.149	91	TTR CEL	-4.985	47.572	-10.098
	91	TAS CES	-0.315	49.421	-17.472	41	718 CZ	-1.794	48.562	-21.463
5	91	TTE DH	-8.182	48.752	-21.744	9.2	ALA M	-4.895	49.958	-14.104
Э	92	ALA CA	-4.547	54.199	-12.747	92	ALA C	-5.823		
	92	ALA D	-6.723	38.176					50.033	-11.903
		_			-13.050	92	ALA CS	-3.197	\$1.621	-12.488
	13	TAL M	-5.959	48.993	-31.329	*3	WAL CA	-7.183	48.834	-10.325
	95	VAL C	-6.708	49.814	-8.899	93	TAL 0	-4.161	47.993	-8.372
	13	WAL CO	-7.957	47.555	-10.411	13	VAL CEI	-9.213	47.488	
	93	TAL CEZ	-8.175	47.376	-12.072	94	LVS B			-9.725
	94	LTS CA	-4.378	50.464				-6.997	50.237	-0.321
		FA2 0	-8.458		-6.999	94	TAR C	-7.331	47.785	-5.874
10	94			30-480	-5.783	94	LYS CB	-4.051	51.976	-6.81 R
	94	LYS CG	-5.394	\$2.320	-5.467	94	L75 CD	-4.848	53.785	-5.502
	94	LTS CE	-4.377	54.208	-4.199	94	LYS EZ	-3.735	33-344	
	95	TAL M	-4.909	49.071	-5.024	93	VAL CA	-7.644		-4.307
	95	VAL C	-4.919	40.491	-2.548				48.457	-3.920
	95	VAL ES				95	VAL D	-7.425	48.154	-1.501
			-1.104	47.030	-4.319	95	ANT CCI	-3.868	44-852	-5.419
	95	TAL CEZ	-6.900	44-180	-4.332	96	LEU G	-5.676	40.974	-2.484
	74	LEU CA	-4.782	49.103	-1.484	94	LEU C	-4.331	58.559	
15	94	LEU O	-3.942	\$1.121	-2.334	94	LEU CO	-3.567		-1.321
	94	LEU CG	-3.573	46.799					48.241	-1.573
	14	LEU CD2			-2.072	94	LEU CD1	-2.207	46.184	-2.163
			-4.489	44.082	-1.845	97	CLY M	-4.324	50.975	-8.484
	17	GLT CA	-3.510	52.307	8.287	97	GLT C	-2.363	52.437	0.315
	9.7	ELT O	-1.619	51.443	0.145	98	ALA W	-1.954	53.448	0.758
	78	ALA ES	-0.428	\$5.478	1.510	71	ALA CA	-0.543		
	9.0	ALA C	0.142	53.116	3.917	11	ALA D		54.048	8.945
	• • •	ASP M	-0.504	52.573	2.912			1.313	52.021	1.663
20	• •	ASP BD1				91	TZ BDS	-2.631	\$1.042	6.151
			-2.730	\$0.902	4.883	11	ASP CE	-2.063	\$1.131	5.040
	99	ASP CO	-8.648	51.693	5.175	99	ASP CA	8.101	\$1.418	3.855
	**	ASP C	0.146	\$0.145	3.320	**	ASP G	0.735	49.313	4.029
	100	SLT M	-0.424	49.883	2.168	100	GLY CA			
	100	GLY C	-1.529	47-451	2.00Z			-0.343	48.521	1.615
	101	SER W	-2.342			100	GLY 0	-1.649	44.512	1.479
				41.128	2.908	201	SER CA	-3.542	47.381	3.315
	101	SER C	~4.759	47.894	2.532	101	SER O	-4.752	48.972	1.907
25	101	866 CB	-3.714	47.447	4.817	101	SER OC	-4.411	48.434	5.209
	105	SLT B	-5.821	47.892	2.577	102	SLT CA	-7.8T7		
	102	GLT C	-8.166	44.536	2.528	182	SLY 0		47.422	1.894
	103	6LW 0	-9.377	47.858	2.478			-7.988	45.431	3.030
	103	SLH C	-10.763			103	GLM CA	-10.535	44.297	3.020
				45.232	2.022	103	GLB	-10.779	45.482	0.017
	103	Pru co	-11.671	47.307	3.274	103	ELM CG	-11.348	48.005	4.584
	103	ELH CD	-12.340	49.104	4.915	103	6L# 0E1	-12.159	49.814	5.902
	103	GLW WEZ	-13.419	49-197	4.112	184	TYP M	-11.611		_
30	184	TYR CA	-12.448	43.124	1.584	194	Tre C		44.141	2.451
	184	TTE D	-12.939	43-276				-13.031	43.470	0.473
	104	TTR C6			-0.487	3.04	TTR CS	-32.497	41.864	2.143
			-11.629	40.829	2.472	104	TTE CD1	-33.019	39.769	3.377
	104	TTE CD2	-10.379	48.753	1.840	104	TTR CEL	-10.005	30.005	3.707
	184	TAB CES	-9.352	49.057	2-171	104	TYR CZ	-9.544		
	104	TTE On	-8.481	30.191	3.324	105	568 B		39.022	3.001
	105	SER CA	-14.877	45-144				-13.909	44.572	4.903
	105	124 4	-14.759		-6.034	105	SER C	-14.172	45.920	-1.159
35	105			43.935	-2.254	105	see ce	-15.880	44.121	0.601
-		SER DC	-15.209	47.834	1.450	106	TEP N	-13.679	44.425	-0.834
	144	TRP CA	-12-421	47.391	-1.948	104	TRP C	-11.895	44.436	-3.012
	104	ter o	-12.021	44.648	-4-2-5	164	TRP CS	-11.321		
	104	TRP CG	-11.643	49.111	-0.204	104	TRP CD1		48.254	-1.355
	104	TEP COZ	-10.658	49.812				-12-162	49.524	8.244
	104				8.501	104	144 261	-12.491	31.358	1.340
		ter ces	-31.351	\$0.573	1.541	104	TEP CES	-9.275	49.852	8.574
	104	TEP CZZ	-10.671	53.318	2.500	106	TPP (1)	-1-441	\$6.56)	1.525
40	184	TOP CHZ	-7.213	51.293	2.455	107	ILE G	-11.339	45-330	-2.481
40	107	ILE CA	-10.765	44.250	-3.325	107	ILE C			
	107	ILE D	-11.675	43.474				-11.955	43.594	-4.190
	107	ILE CAL	-8.634	43.764	-5.398	107	ILE CS	-1.944	43.10)	-2.523
	107				-1.934	107	IFE CCS	-9.632	41.730	-3.361
		ILE COI	-0.213	42.998	-8.627	101	14! #	-12.904	43.292	-3-577

50

	100	ILE CA	-14.114	42.122	-4.373	394	ILF C	-14.639	43.474	-5.386
	100	ile v	-24.874	43.324	-6.552	100	ILE CO	-15,246	42-245	-3.326
		ILE CEI	-14.724	41-677	-2.482	109	ILE CG2	-14.540	42.024	-4.095
	1	116 (01	-15.452	48.845	-1.111	101	45m b	-14.751	44.958	-4.981
	111	450 64	-15.204	46.018	-5.914	109	ASM C	-14.232	46.867	-7.004
	117	ASM 0	-14.440	48.272	-8.235	199	ase ce	-15.208	47.359	-5.207
5	***	ASH CG	-14.578	47.486	-4.353	107	458 831	-17.455	46.675	
•	167	438 802	-14.633	48.447	-3.442	110	SLT #	-12.951	45.700	-4.646
	110	GLT CA	-11.752	45.917	-7.865	110	617 6	-12.108	44.712	-4-774
	110	617 0	-11.929	44.929	-10.034	111	ILE .	-12.379		-8.612
	111	ILE CA	-12.403	42.334	-9.011	111	ILE C	-13.059	43.539	-8.246
		ILE #	-13.921	42.384	-11.148	111	TLE CO	-12.734	42.540	-9.942
	111	116 (61	-11.421	40.501	-7.655	111	ILE CEZ		48.948	-0.344
	111	116 601	-11.548	31.786	-6.334	312	CLU B	-23.122	39.791	-9.347
10	111		-16.118	43.374	-19.844	112	610 (-14.893	43.875	-9.240
	112	ELU CA					GLU CS	-15.672	44,347	-11.171
	112	ELU O	-14.447	44.130	-12.246	317		-17.229	43.099	-9.141
	112	ern ce	-17.847	42.917	-8.135	112	ELU CO	-14.724	41.024	-8.485
	315	CLU DET	-19.941	40.844	-0.014	112	ern ses	-19-123	41.921	-9.844
	113	TEP &	-15.014	45.403	-10.971	113	TEP CA	-14.754	44.408	-12.000
	113	TRP C	-14.876	45.443	-13.140	313	TRP D	-14.319	45.932	-14.332
	113	TRP CO	-13.002	47.553	-11-434	113	TAP CE	-13.486	48.334	-12.401
15	333	TRP CD1	-14.148	49.736	-12.681	113	TRP CD2	-12.441	40.552	-13.463
	213	TRP WEL	-13.597	\$8.443	-13.723	113	TRP CEZ	-12.545	49.741	-14.215
	113	Tep (63	-11.451	47.445	-13.809	113	TRP CZ2	-11.676	50.845	-15.274
	113	TRP C23	-10.410	47.199	-14.879	113	TRP CH2	-10.752	49.874	-15.603
	314	ALA M	-13.019	44.801	-12.632	114	ALA CA	-12.333	44.045	-13.874
	314	ALA C	-13.199	43.179	-14.752	114	ALA O	-12.943	43.674	-15.978
	114	ALA CB	-11.299	43.192	-13.140	115	ILF W	-14.174	42.540	-14.119
	115	ILE CA	-15.070	41.640	-14.097	115	ILE C	-11.920	42.485	-15.056
20	113	ILE D	-34.977	42.225	-17.070	115	ILE CO	-16.000	40.840	-13.922
	113	ILE CEL	-15.218	39.036	-13.043	115	IFE CES	-17.151	40.168	-14.755
	115	ILE COL	-14.004	39.411	-11.743	114	ALA N	-14.534	43.527	-15.207
	314	ALA CA	-17.390	64.448	-14.050	114	ALA C	-14.764	45.047	-17.278
	114	ALA D	-17.323	45.255	-18.343	116	ALA CB	-18.011	45.510	-15.151
	117	ASK H	-15.423	45.390	-17.122	117	ASH CA	-14.553	45.947	-18.139
	117	ASM CO	-13.427 -13.615	44.974	-14.034	117 117	ASM CE	-12.997	45.436	-19.820
25	117	ASM DD1	-14.565	44.958	-17.424	117	ASH HDZ	-14.400	48-177	-16.939
25	114	ASM M	-14.223	49.612	-17.773 -18.967	114	ASH CA	-14.931	48.249	-15.736
	111	ASM C	-12.240	42.444	-19.943	110	ASH D	-13.760	42.642	-19.032
	111	ASH CB	-14.247	42.843	-21.279	118	ASH CE	-11-617	42.309	-20.932
	111	41% OD1	-16.510	42.321	-20.759	110	45H MO2	-15.737 -16.136	43.040	-21.395
	111	MET W	-11.486	42.500	-18-475	117	MET CO	-10.232	44.016	-22.133
	111	967 6	-10.025	48.734	-18.928	119	RET O	-10.888	42.222	-18.478
	111	MET CB	-9.410	42.461	-17.055	219	MET CG	-9.880	39.638	-18.759
30	317	MET SO	-8.788	44.943	-17.526	119	RET CS	-9.982	43.883 44.861	-16.582
	120	ASP W	-8.904	40.437	-17.584		ASP CA	-1.410		-18.763 -20.830
	120	ASP C	-7.822	34.310	-10.856	120	ASP G	-1.0)8	39.116 37.107	
	120	ASP CB	-7.555	37-154	-21.234	120	43P E6	-8.237	39.736	-10.690 -22.454
	120	ASP 001	-7.001	40.704	-23.884	150	859 002	-9.327	37.135	-22.139
	121	VAL W	-7.021	39.117	-18.115	121	TAL CA			
	121	VAL C	-6.296	39.534	-15.706	121	VAL D	-6.226 -6.284	38.681 40.788	-14.974 -15.909
_	121	VAL CO	-4.735	38.587	-17.494	121	TAL CGS	-3.758	38.174	-10.427
35	121	VAL CEZ	-4.787	37.916	-18.844	155	ILE a	-4.310	38.976	-14.570
	122	ILE CA	-4.248	39.797	-13.397	122	ILE C	-5.020	39.242	-12.427
	122	216 6	-4.829	38.012	-12.469	322	Ire ce	-7.476	37.604	
	122	ILE CGI	-8.484	48.392	-13.043	122	ILE CES	-7.221	39.663	-12.466 -18.954
	122	FLE COI	-9.974	39.784	-12.31)	173	459 9	-4.263	40.222	-12.114
	153	ASW CA	-3.145	39.854	-11-232	12)	ASM C	-3.562	40.404	-9.841
	123	410 8	-3.708	41.631	-9.833	173	ASH CS	-1.828	48.478	-11.497
40	123	450 (6	-0.492	40.048	-10.777	123	450 001	~8.04)	30.770	-11.010
40	123	41E 002	-0.346	40.747	-9.720	124	att m	-3.450	39.404	-8.632
	124	HET CA	-1.410	19.073	-7-414	124	#17 F	-2.471	30.403	-0.414

		4	-1 50.		-4.813	124 881 68	-4.943	34.317	-4.171
	124	#E7 0	-3.304	31.500				30.472	
	11.	et 1 £6	-6.198	48.112	+7.473	124 - 47 82	-7.586		-4.150
	114	417 E1	-7.948	31.075	-7.542	125 680 w	-1.414	40.491	-4.50}
	111	814 64	-0.193	48.287	-3.761	123 870 6	-8.622	48.712	-4.314
	121	\$11 0	0.233	41.617	-3.003	125 BT4 C9	1.021	41.027	-4.321
	111	344 86	1.444	40.494	-7.875	124 LTU M	-1.431	48.678	-8.778
5	124	LEU CA	-1.442	40.347	-2.304	174 LEU C	-2.438	21.214	-1.807
3	12.	110 8	-1.0.	38.134	-2.129	126 LEU EB	-2.791	41.548	-2.410
							-5.278	41.131	-2.574
	116	FER CC	-3.911	41.447	-3.333				
	114	FEN CBS	-6.179	42.740	-4.073	127 GLY W	-2.522	39.412	-1,411
	111	BLT CA	-3.835	37.071	0.193	127 BLT C	-3.174	38.380	2.482
	117	647 0	-2.446	34.010	2.220	111 GLT B	-4.121	37,443	2.212
	111	GLT CA	-4.415	37.486	3.442	229 BLY C	-4.644	36.938	4.104
	121	SLT B	-4.983	35.154	3.274	129 000 4	-4.519	35.657	8.402
10	111	PRE EA	-0.671	34.323	8.991	129 P40 C	-4.116	34.884	4.482
, 0	121	PIC D	-4.334	32.117	4.305	111 015 61	-4.040	34.484	7.314
					7.727	119 P05 CD	-4.231	34.870	4.414
	121	P10 C6	-4.419	34.314			-8.670	34.411	4.023
	130	111 W	-7-051	33.013	6.932	130 Sta Co			
	110	810 6	-9.218	34.884	4.726	130 117 3	-1.141	31.681	4.029
	130	811 68	-9.549	35.351	7.210	130 \$28 06	-1.123	34.424	4.493
	131	GLT N	-10.003	33.947	4.141	131 BLY C#	-10.624	\$4.229	3.674
	131	SLT C	-12.205	34.713	1.542	131 GLT D	-12.495	34.722	4.751
15	132	\$ E B N	-11.940	35.011	2.574	112 88* 64	-14.407	35.433	3.011
	132	\$14 E	-15.289	34.805	1.134	172 862 0	~14.791	24.514	8.824
	132	siè te	-10.590	34.927	3.141	132 112 06	-14.493	27.537	1.175
			-14.547	14.541	2.204	173 ALA CA	-17.507	34.037	1.324
	1))	ALA R					-17.741	34.437	-1.014
	133	ALA C	-17.480	34.745	0.007	133 ALA D			
	133	ALA EB	-14.866	83.628	1.116	134 ALA W	-17.683	34.241	0.254
	134	ALA CA	-17.672	87.259	-0.792	134 ALA C	-14.435	37.369	-1.674
	134	ALA D	-14.781	37.585	-2.747	134 ALA CB	-16.243	38.400	-8.197
20	133	LEU M	-15.478	17.229	-1.046	133 LEU CA	-14.197	37.244	-1.604
	133	LEU C	-14.130	36.005	-2.761	138 LEU 0	-11.794	36.825	-3.410
	135	LEU EB	-13.038	27.320	-0.718	135 LEU CE	-11.493	37.130	-1.500
	135	LEU COI	-11.460	30.415	-2.212	128 640 601	-10.582	34.807	-1.119
	136	L75 &	-14.909	8435	-2.173	136 LTS CA	-14.543	33.517	-3.011
	114	LTS E	-13.544	13.739	-4.110	136 175 0	-19.279	21.431	-0.305
	134	LTS CB			-2.184	136 LV1 CG	-14.743	31.047	
			-14.901	32-341					-3.643
	3.34	LTS CD	-15.01)	20.472	-2.134	336 L75 CE	-15.741	23.707	-2.774
25	134	FAR #1	-15.301	28.411	-4.140	337 ALA W	-16.704	34.240	-3.947
	137	BLB CB	-17.795	34.414	-4.813	137 ALA C	-17.338	38.303	-4.443
	137	ALA D	-17.709	31.049	-7.208	137 ALA EB	-19.094	34.743	-4.243
	111	ALA M	-16.529	84.301	-3.729	138 ALA CA	-14.901	37.311	-4.411
	111	BLA C	-14.953	24.474	-7.557	178 AL* D	-14.985	26.843	-8.762
	331	ALA CS	-15.522	24.547	-5.934	139 VAL &	-13.950	33.759	-7.827
	111	VAL CA	-12.944	89.291	-7.837	139 V41 C	-13.423	34.224	-8.720
	111	TAL D	-13.206	34.070	-9.877	130 VAL CO	-11.830	34.673	-4.761
30	130	VAL CG1	-10.010	83.854	-7.844	139 VAL CG2	-11.078	31.780	-6.213
50	140	ASP N	-14.993			140 45" [8		32.496	
				33.134	-8.123		-28.274		-8.121
	1.0	ASP C	-14.823	33-131	-10.044	1.0 45.0	-14.000	37.579	-11.198
	140	ASP CB	-14.149	31.549	-1.111	347 459 66	-25.388	38.640	-7.184
	140	41 P 001	-14.178	30.403	-7.212	1+0 ASP DTZ	-14.139	30.132	-4.319
	141	LTS W	-14.451	34.263	-9.810	141 E71 CA	-17.373	38.004	-14.568
	141	LTS C	-14.373	35.415	-12.746	143 L73 D	-10.700	31.240	-13.111
	141	LTS CO	-11.939	36.275	-10.311	141 LYS CG	-16.884	37.014	-11.306
35	141	LTS CO	-11.594	24.187	-19.531	141 LYS CE	-28.572	34.051	-11.250
•	141	LTS AT	-21.136	40.037	-10.273	142 414 4	-15.167	21.047	-11.544
	1.1	ALA EA	-14.173	36.192	-12.414	1+2 414 6	-13.010	35.010	-13.521
			-13.770					34.697	
	141	414 0		35.169	-14.755		-12.870	32.705	-11.941
	143	TAL N	-13.912	33.114	-12.432	343 AVT CV	-13.148		-13.450
	143	VAL C	-14.346	33.331	-14.476	143 VAL C	-10.100	31.184	-15.619
	143	VAL EB	-12.551	31.473	-12.714	143 VAL CG1	-12.100	38.370	-13.441
	141	ANT CES	-11.101	32.191	-12.014	344 ALA N	-11.581	32.238	-13.875
40		ALA FA	-14.844	81.884	~ 1 4 - 3 4 1	SAA ALA P	-14.914	81.401	-18.841

	_			32.243	-14.953	144 414 58	-17.942	31.941	-13.780
	244	AL4 E	-11.380		-13.764	145 \$19 CO	-16.682	84.917	-14.784
	341	311 h	-16.507	33.9.8					
	141	3 * * 6	-11.609	34.773	-17.019	14) \$ER D	-11.910	33.321	-18.893
	141	110 60	-17.010	34.374	-14.614	148 889 00	-11.533	36.931	-19.849
	144	GLT N	-14.877	33.414	-17.545	144 BL7 E4	-13.619	33.741	-14.675
				24.491	-12.385	144 BLY D	+11.420	24.344	-19.266
5	146	er. c	-11.173			147 VAL CA	-10.074	88.884	-14.911
٠	147	TAL R	-12-170	35.112	-17.254		-10.171	30.991	-15.484
	141	ANT E	-4.830	34.634	-14.323	144 ANT D			
	147	TAL CB	-11.152	36.977	-15.004	147 WAL CG1	-9.894	37.803	-15.575
	1.7	WAL EES	-12.340	37.935	-14.230	345 WAL W	-8.513	36.014	-14.663
	-	VAL CA	-7.482	34.230	-14.808	148 VAL C	-7.187	34.907	-14.701
	141				-14.750	148 VAL CS	-4.173	34.114	-14.930
	141	VAL D	-4.845	24.133			-4.390	33.432	-18.262
	148	WAL EGS	-5.079	33.483	-14.241	Jes Auf Ees			
	144	TAL R	-7.251	34.385	-13.531	347 VAL CA	-4.117	34.945	-12.249
10	147	WAL C	-6.700	34.381	-11.613	149 VAL 0	-5.624	32.173	-11.439
	149	VAL EB	-6.224	34.890	-11.313	149 VAL CEI	-7.893	38.419	-11.001
	-				-12.094	150 VAL W	-4.732	35.301	-11.404
	149	ANT CES	-1.414	31.346			-3-157	35.621	-9.557
	110	WAL CA	-3.563	34.987	-10.901				
	100	WAL D	-3.512	34.778	-9.400	190 VAL CB	-2.274	35.343	-11.991
	150	TAL EGS	-0.973	34.433	-11.461	150 WAL CE?	-2.475	34.943	-13.361
	111	44.4	-2.548	34,744	-8.315	151 ALS CS	-2.341	30.342	-7.287
					-4.457	151 464 0	-0.616	23.811	-4.984
15	191	ALA C	-1.000	39.034			-0.410	35.907	-3.922
	151	ALA CO	-3.557	35.340	-6.307	155 ALA W			
	152	ALA EA	0.714	21.438	-5.117	187 ALA C	0.304	34.310	-4.161
	111	ALA D	-8.728	34.464	-3.447	182 ALA CT	1.246	34.697	-4.294
	193	ALA M	1.125	33.102	-3.912	193 ALA CA	8.840	32.256	-2.943
		ALA C	0.931	32.725	-1.511	153 ALA 0	8.317	32.392	-0.541
	111					154 BLY 4	1.627	22.442	-1.244
	111	ALA CO	1.750	21.030	-3.195				
	154	ELT CA	2.043	84.233	0.123	184 BLT C	3.519	24.040	0.330
	354	SLT D	4.100	13.267	-8.118	195 ASW N	3.711	34.788	1.560
20	111	BEN CA	5.344	34.787	1.037	133 454 C	3.111	34.250	3.442
	111	ALV P	4.101	84.429	4.293	185 ASM C9	4.008	34.170	1.964
					0.700	155 ASN 001	4.123	34.045	-0.514
	135	484 CG	3.210	86.702			4.711	22.161	1.475
	193	ash md2	1.434	27.945	0.352	154 BLU b			
	156	GLU CA	4.433	82.537	4.970	194 ELU C	9.522	31.328	9.107
	156	ELU D	9.374	30.437	4.222	186 GLU CD	3.103	31.900	5.100
	114	SLU CG	2.471	32-442	4.348	190 ELU CO	2.114	33.911	4.278
	134	SLU DES	1.744	14.322	5.312	194 GLU DE2	3.104	34.456	7.146
25					4.227	197 BLT CA	7.104	20.917	4.387
	257	SLT N	6.311	31.057			5. 114	28.344	4.001
	257	GLT C	4.507	20.622	4.313	157 GLT D			
	111	TAR W	7.147	27.793	3.312	398 THP CG2	8.079	21.374	3.850
	116	THE DEL	8.707	25.487	4.217	198 THP CB	7.864	25.344	8.294
	111	THE CA	4.552	20.467	9.702	181 THE C	6.190	24.480	7.197
	111	THE D	4.479	27.335	7.977	199 114 4	8.331	28.441	7.497
						119 668 60	3.473	24.105	9.212
	111	26. 00	3.141	25.904	30.315		4.494		4. 544
30	111	SIT CA	4. 831	25.210	1.111	355 368 6		23.720	
30	111	511 D	3.339	23.261	9.835	240 BLY M	3.574	22.947	8-831
	100	BLY CA	5.434	21.504	4.415	240 BLY C	4.570	22.449	7.734
	140	BLY B	4.608	21.324	4.313	141 St. W	3.425	20.810	6.114
				19.777	7.014	141 114 6	1.477	26.706	4.784
	161	\$ E + C A	4.414		9.841	341 888 68	2.344	18.173	7.271
	161	814 0	8.494	20.347					7.451
	161	811 00	1.114	18.028	4.515	167 188 4	1.103	81.841	
	162	888 64	0.167	22.721	7.113	185 PE. C	0.430	23.112	8.044
	142	311 0	1.533	23.040	3.394	102 584 69	-6.213	23.444	8.241
35	167	310 06	8.194	23,091	9.486	111 111 6	-8.479	23.921	8.197
				24.750	1.912	24) 188 6	-8.441	24.177	4.513
	143	810 E4	-8.411				-1.000	24.442	3.211
	143	864 D	-1.070	86.141	3.504	141 SEC CO			
	143	880 00	-1.992	83.718	3 - 333	304 TPR W	9.357	26.912	3.637
	144	THE CA		29.340	4.312	164 THR C	8.109	29.284	3.194
	144	141 0	1,413	30.502	3.270	166 THE ES	2.015	20.510	4.814
	144	7#P 061	1.114	26.202	3.492	104 THE EG2	3.317	27.410	4.001
			-0.313	28.742	2.190	141 VAL CA	-0.919	19.1-2	1.014
40	103	VAL N			1.097	141 TAL D	-2.020	30.132	1.200
. •	3.65	WAL C	-2.024	30.541		141 445 0	-8.747	*****	

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	141	WAL CO	-1.111	21.124	-8.161	145 WAL CS1	-1.947	29.351	
	101	TAL ES?	-3.216	27.714	-0.111	166 617 4	-1.918		-1.114
	1	BLT CA	-2.743					31.021	1.129
				32.776	1.616	\$44 ELT C	-4.818	32.891	0.617
	160	GLY D	-4-124	32.184	-8.316	167 770 4	-5.614	33.731	4.979
_	867	118 CO	-4.823	34.144	0.113	167 TTP C	-1.913	35.289	-8.686
5	167	111 0	-1.676	36.213	8.114	347 778 68	+7.464	34.232	
	167	111 66	-7.791	37.114	1.709	167 Tra CD1	-7.204		1.244
	167	TTE CD2	-8.710					32.703	2.947
				32-114	1.133	367 778 621	-7.567	31.524	3.418
	167	111 (12	-9.941	30.011	3.804	167 TTE CI	-8.484	30.671	3.044
	167	711 D-	-6.816	24.483	3.451	166 PRD N	-6.310	31.499	-1.850
	340	P16 66	-4.943	34.374	-3.931	168 P#0 CD	-4.273	34.782	-1.614
	166	PRC CO	-7.804	35.344	-3.505	168 *05 CA	-7.134		
	166	P40 C	-4.311	33.134	-3.270			34,457	-2.860
10	149	6L7 M					-7.097	35-850	-3.712
			-5.000	33.193	-3.109	169 ELT CA	-4.444	32.677	-3.927
	100	SLT C	-4.937	30.701	-3.470	169 BL7 D	-4.800	29.733	-4.249
	170	F42 #	-5.402	30.179	-2.255	378 L73 CA	-3.814	27.265	-1.748
	170	611 6	-7.055	28.773	-2.514	370 175 0	-7.301	27.854	
	170	LTS CB	-4.244	29.214	-0.284	170 LV1 CG	-5.795		-2.524
	170	LTS CO	-6.250	21.211				20.106	8.551
					2.031	370 LTS CE	-5.733	27.271	3. 629
	170	FAR WE	-4.259	27.413	3.215	372 TTE N	-7.832	29.116	-3.141
15	171	44 CA	-9.912	29.043	-3.837	171 TTD C	-8.683	24.101	-5.113
	171	111 D	-7.760	28.714	-8.928	171 TYR CO	-9.942	30.224	-4.242
	171	778 E6	-10.497	30.484	-3.047	171 TTR CD1	-11.940		
	171	TTR CD2	-10.454	32.374	-2.026			30-303	-1.962
	171	110 CE2	-10.941			171 178 CE1	-11.510	31.003	-8.867
				33.063	-3.936	171 TTR C2	-11.520	32.311	-3.816
	171	111 0-	-11-808	33.119	8.170	172 P#0 m	-1.217	27.204	-3.274
	171	PAC CA	-9.013	24.417	-4.344	172 PRO C	-9.233	27-154	-7.669
	172	720 D	-0.525	24.784	-0.681	272 PRD CB	-10.167	25.329	-4.513
20	172	P40 C5	-18.600	29.271	-8.056	372 080 CD	-10.364	24.441	-4.11.
	171	511 4	-10.057	28.167	-8.019	173 158 CA	-10.220		
	175	111 (-9.025	29.773				25.118	-0.330
	171	111 61			-9.591	173 11° D	-1.744	30.233	-18.742
			-11.528	21.623	-9.481	178 SEP DG	-11.595	30.544	-8.404
	374	TAL M	-0.142	29.944	-8.414	374 VAL CA	-7.852	30-091	-1.155
	374	VAL C	-5.754	30.131	-9.068	874 VAL D	-5.612	21.132	-1.344
	174	VAL CB	-4.271	31.775	-7.594	STA VAL CG1	-5.714	\$2.837	-7.617
	274	ANT CES	-6.220	32.503	-1.323	LTS ILE W	-4.911	\$0.729	-0.00
25	175	ILE CA	-3.849	36.156	-10.024	179 1LE C	-2.714		
	111	111 0	-2.450	31.958	-8.955	ifi ice ca		30.734	-8.694
	175	1LE C61	-3.857	29. 174	-12.524		-2.933	30.524	-11.41*
	175	ILE CEL				175 1LE CG2	-1.451	30.019	-11.512
			-3.692	30.529	-13.944	174 dia w	-1.219	30.028	-7.925
	176	ALA CA	-1.311	30.517	-6.870	274 ALA C	8.120	30.303	-7.310
	174	ALA D	4.433	29.215	-7.838	176 BLA EB	-1.419	27.116	-3.541
	177	TAL B	8.844	31.410	-7.180	177 VAL EA	2.241	11.534	-7.454
	177	VAL C	3.223	31.493	-4.473	177 VAL D	3.178		
30	277	MAL ED	2.431	32.607	-8.765			32.457	-8.721
	117	VAL CEZ	1.374	32.552	-1.141		3.462	32.447	-1.312
	170	SLY CA				178 BLY N	4.877	30.654	-4.398
			1.141	30.703	-5.339	170 BLT C	6.444	31.233	-6.874
	178	6L1 0	4.491	31.436	-7.216	179 ALA W	7. 912	31.447	-1.267
	179	ALA CA	0.715	32.037	-3.851	170 ALA C	9.939	31.074	-3.771
	179	ALA C	10.198	30.401	-4.719	179 AL4 CB	9.025	33.211	-4.973
	180	TAL &	10.410	31.142	-4.005	180 VAL CA	11.970		
	100	VAL C	11.046	33.515	-7.171			30.412	-4.981
35	145	VAL CO	12.075			180 VAL B	12.712	32.411	-7.417
				20.514	-8.144	180 ANT ERT	11.271	24.251	-7.811
	180	ANT EES	11.675	30.110	-9.500	18: ASP M	84.267	31.201	-6.810
	111	ASP CA	13.433	32.200	-7.011	101 ASP C	11.942	31.804	-6.442
	\$ 8 1	417 0	11.339	31.890	-9.212	101 410 62	14.444	81.921	-5.014
	101	ASP-CG	17.120	30.534	-5.971	181 ASP 001	17.103	20.713	
	181	43P 002	17.680	30.214	-4.887	302 800 6			-6.972
	112	311 64	17.622				17.067	32.384	-9.847
	111	\$1 D		37.214	-10.171	101 BFF C	11.177	30.017	-20.494
40			16.365	30.432	-11.670	193 864 66	48.478	33.313	-15.444
	185	81+ D6	48.014	34.561	-10.475	183 \$f* m	10.255	30.942	-9.423
	10)	880 -Ca	18.716	28.645	-9.444	18) 810 C	17.511	27.614	-9.547
	111	84 0	17.859	20.415	-9.397	383 384 60	19.236	20.323	-8.997
						, us			

	103	11 1 86	23.565	20.615	~0.251	104	ASA W	14.373	28.004	-1.411
	104	454 64	15.144	27.317	-9.400	104	Alm t	\$4.971	24.720	-8.197
	164	AT - D	14.136	25.769	-0.097		454 CB	\$8.914	24.341	-16.722
	10.	41 4 66	14.993	24.991	-12.074	184	414 831	14.780	28.184	+11.277
	184		11.11:	20.210	-11.074	101	SLR &	15.342	27.247	-7.159
5								14.290	27.494	
a	113	BL# CA	11.274	24.444	-5.833		GIN C			-1.26)
	103	EL* D	14.159	28.724	-1.114	100	GLW (B	14.979	24.540	-3.101
	183	SLA CC	14.119	24.242	-3.614	188	SL= CD	10.011	26.182	-3.294
									24.384	-1.934
	141	61 = 013	10.164	25.700	-4.141		gra ats	11.244		
	194	416 4	13.278	24.411	-4.448	114	43 384	12.183	21.774	-2.841
	16.	486 C	12.780	28.762	-2.844	111	406 0	13.678	20.384	-2.017
		416 64	11.111	20.143	-3.110		496 EG	10.214	27.471	-2.141
	3 6 4									
	18.	ARC CD	1.447	24.137	-1.448		416 48	4. 4.4	24.333	-0.117
10	104	ARG CI	9.661	24.471	1.000	100	436 mm1	9.347	27.880	1.65#
	111	686 442	10,946	24.321	1,713		ALA N	12.294	30.011	-2.813
	387	ALA CA	32.726	31.004	-1.893		ALA C	12.262	30.404	-6.817
	157	ALS D	11.198	30.043	-0.317	187	ALA CE	11.144	32.403	-2.344
	111	511 -	13.001	36.770	0.547	100	580 CA	22.671	30.204	1.661
							510 0		30-111	3.211
	100	314 6	11.314	30.847	2.412			38.740		
	100	811 68	23.747	30.414	2.931	168	88. 00	84.137	31.034	Z. 041
	100	PHE W	10.143	32.010	1.974	141	PHE EA	9.697	32.681	2.418
15	111	P=1 6	4.474		1.609		PHE D	7.347	32.014	2.011
, 0				\$2.194						
	199	*#[CB	9.787	34.217	2.243		PAT CG	10.317	34.498	8.867
	141	PHE COL	9.147	34.130	-1.121	189	946 CD3	11.419	30.114	0.547
	111	PRE CAS	9.483	33.117	-1.411		PHE CEZ	11.769	30.141	-6.761
	300	PRE CI	19.786	35.516	-1.725		SEC N	8.703	31.524	0.411
	840	BER CA	7.626	31.094	-0.391	190 1	884 6	6.843	30.162	0.321
	190	84 0	7.834	27.013	8.844	190	510 CB	8.181	30.390	-1.788
	100	811 06	7.186	30.387	-2.618		\$ E # #	1.111	10.771	0.214
20										
20	291	311 CA	4.341	27.674	8.987		81+ C	4.261	28.330	9.223
	191	141 0	4.843	24.269	-0.175	391	584 EB	3.015	30.411	0.913
	191	511 DG	2.720	31.245	1.954	392	VAL W	3.756	27.310	0.924
	105	TAL CA	3.421	23.612	0.391		ANT C	2.254	23.291	0.414
	7 . 5	VAL D	1,851	25.610	1.198	192	ATF CO	4.781	20.127	1.911
	192	WAL CEI	8.144	23.727	0.722	192	TAL CGR	4.417	26.104	2.512
	193	GLT N	1.111	24.172	0.047		BLT CA	0.629	13.514	0.415
	173	SLT E	0.081	23.029	-8.901		8L4 D	9.530	23.244	-2.615
25	194	PIC W	-1.023	22.289	-0.722	294	PED CA	-1.462	21.001	-1.873
	114	PRE C	-2.237	12.605	-2.614	194	P 6 D . D	-2.41)	22.244	-4, 915
	194	PRD 68	-2.749	20.763	-1.210		PRD 66	-2.311	20.622	0.213
	194	POD CD	-1.633	21.754	8.171	195	BLU N	-2.122	23.793	-2.431
	111	SLU CE	-3.148	24.850	-3.252	193	BLU C	-2.015	20.411	-4.051
	101	SLU B	-2.518	24.371	-4.114		BLU CO	-4. 643	29.784	-1.478
		: .								
	175	ern ce	-4.742	25.124	-1.475		Prn CD	-4.715	24.860	-0.100
	195	BLU DEI	-3.110	24.960	D.145	198 (ELU 881	+5.130	24.520	•,703
30	194	LEU N	-0.124	25.264	-3.870	194	LEU CA	8.241	20.921	-4.444
							LEU C	8.308	14.111	-6.113
	194	LEU E	0.121	21.374	-6.039					
	196	LEU CE	1.340	25.789	-3.654	294	LEU CE	8.770	24.170	-4.643
	194	LEU EDA	2.730	27.714	-4.631	196	LEU CDI	4.627	25.721	-3.911
	197	417 1	4.140	26.208	-7.013		417 64	0.032	28.774	-8.488
	197	437 2	1.307	31.731	-1.211		41° 0	1.455	24.734	-9.914 -
	197	45" (8	-1.047	24.511	-9.191	197	ASP EG	-2.404	14.351	-8.541
	197	ASP 001	-1.804	29.155	-1.314		417 002	-3.035	27.317	-1.011
35										
	290	VAL N	2.013	24.011	-7.3.4		ANT EV	3.204	26.970	-18.200
	198	VAL C	4.157	27.930	-9.514	190	ANT D	3.752	28.011	-8.587
	194	VAL CA	2.414	27.474	-11.637		VAL CG1	1.930	26.724	-12.937
	194	TAL CER			-11.414		MET W	5.374	27.911	-18.616
			1.337	31.919						
	300	MET CA	4.431	28.802	-9.498		MEA C	4.843	29.010	-18.578
	399	# 21 D	4.474	25.316	-11.777	198	MET CS	7.660	27.970	-9.877
	199	#87 CG	7.343	24.949	-8.139		BTT 10	4.753	27.441	-4.541
					-1.117		ALA A	1.424	30.942	-10.163
40	111	MET CE	1.227	27.735						
40	200	BLA CA	7.771	31.929	-11.015		ALA C	1.111	11.040	-18.272
	200	ALA D	0.127	32.924	-9.840	200	ALA CB	4.932	32.078	-21.416
									_	-

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				9. 921	32.499	-10.911		PRE C4	11.013	\$4.130	-10.231
	713						201				
	201	,,,		10.430	39.127	-9.231	201	981 9	0.579	33.907	-4.482
	111	710 6		81.017	34.722	-11.400	801	33 344	11.393	14.040	-18.678
	201	440 C		8.841	33.414	-12.405	101	SLT R	10.928	30.204	-8.821
	111	BLT C		10.473	34.234	-1,9-4	202	BLY C	11.880	34.471	-4.111
	202	BLT E)	11.352	37.124	-4.575	103	VAL B	12.015	84.513	-6.613
_	203			11.941	34.529	-1.714	203	WAL &	14.786	38.017	-6.441
5	103	VAL E		15.133	37.731	-7.593	101	VAL ER	14.814	11.411	-5.361
	10)			14.974	34.104	-4.412	103	V41 662	14.179	34.741	
											-4.375
	104	111		14.003	34.182	-3.839	204	\$20 C4	16.672	48.231	-6.487
	204	311 6		18.047	40.619	-7.872	104	804 C	18.784	49.481	-1.411
	21.	161 2		17.017	39.974	-8.374	\$04	86. 02	37.732	41.184	-4.472
	101	1.1	1	13.772	45.943	-8.008	201	ILE CO	13.869	41.234	-0.235
	201	111 5		13.207	42.749	-8.478	201	314 0	12.675	43.490	-0.440
10	205	111 0		11.532	40.811	-9.144	205	TLE CES	11.434	31.314	-8.810
10	203	THE C	61	10.277	41.281	-10.467	203	TLE CD1	12.257	30.412	-9.771
	104	6.4 4		13.954	41.015	-10.489	204	SLW CO	14.20.	44.517	-10.634
	264			11.062	44.974	-11.430	206	5L# 0	12.447	44.318	-12.621
	204	-		15.415							
					44.708	-11.740	204	BLN EG	34-684	44.161	-10.980
	\$54	SUN E		17.285	48.1+8	-30.007	204	Erm DEI	10.310	44.934	-9.333
	107	CL N		14.554	46.260	-9.857	201	8 E 4 M	12.319	44.844	-11.214
	101	360		31.217	46.571	-11.987	307	11+ C	11.017	40.013	-11.749
15	267	36.0)	11.419	48.457	-11.004	201	311 61	9.918	45.833	-11.541
	207	\$11 6	36	1.993	46.016	-12.611	201	THE .	10.814	48.664	-12.324
	201	THE E	6.2	9.171	\$0.339	-14.754	100	THP D61	7.870	41.414	-13.144
	101	THE C	4	0.620	\$0.415	-13.357	101	THE ER	9.478	\$0.092	-12.173
	201	1=1 6		8-197	80.488	-10.403	101	THE D	1.423	49.807	-18.649
	201	140		8.454		-10.228	101	LEU EI	9.192	52.250	-4.951
	101	LIU		8.673	\$1.413						
					\$3.410	-1.202	201	LEU D	9.140	54.227	-10.222
	201	LEU C		10.333	52.192	-7.911	201	TER CE	10.804	88.014	-7.414
20	304	FER (21-948	\$1.114	-4.472	101	TEN CBS	9.407	90.202	-4.449
	21¢	PED 4		7.796	94.139	-4.444	810	PRD CA	7.273	\$\$.\$17	-8.441
	210	P#0 (:	6.383	86.573	-8.431	210	PRC G	9.491	54.445	-8.184
	210	**> (6.302	\$5.733	-7.517	210	22 344	4.004	54.379	-4.944
	210	P80 (. 5	7.193	\$3.491	-7.271	211	SLY N	6.977	57-445	-9.333
	211	SLY C	4	9.049	38.743	-9.410	211	BLY E	14.894	51.454	-18.493
	211	647 6		11.176	89.005	-10.259	111	454 4	9.453	37.770	-11.567
	212	ASH I		10.903	\$7.422	-12.643	212	ASh E	12.019	56.753	-12.004
	117	454		17.100	87.161	-12.020	112	451. CO	11.224	\$0.555	-13.499
25	111	414		11.001	59.115	-14.814	212	884 831	11.053	57.654	-15.323
	212	410		32.273	\$1.157	-11.374	iii	LTS N	11.00)	99.749	-11.247
	113	L78 (12.810							
					\$4.746	-10.937	213	LTS E	12.448	\$3.419	-18.866
	211	L73 (31-775	53.039	-21.617	213	LTS CI	12.749	33.241	-9.859
	211	173		13-204	34.474	-8.767	211	LTS CO	13.2.4	37.030	-7.312
	211	171 (14-125	50.210	-4.870	111	F41 85	15.040	88.761	-7.921
	814	178		33.611	32.761	-28.444	214	778 CA	13.802	81.244	-10.722
30	814	148 (24.383	60.600	-1.481	214	777 8	10.211	51.213	-4.817
	214	778 (34.641	80.981	-11.984	214	TTR CS	24.110	91.421	-13.244
	214	778 (: 51	14.619	82.847	-13.478	214	TTR CC2	13.129	51.043	-14.614
	214	774 (i i	14.230	83.475	-14.814	214	TYR CRE	12.45	\$1.649	-15.178
	214	111	1	11.204	\$2.193	-15.110	214	TTD CH	12.754	13.411	-14.696
	211	SLT I		14.858	49.147	-9.154	111	BLY CA	14.422	48.772	-7.903
	215	617		14.111	47.325	-7.749	215	617 0	13.249	46.917	-8.021
	210			14.010	44.616	-4.031	214	ALA CA	14.454	45.203	-4.741
~	216	ALA I		13.402							
35 /					44-922	-1.112	214	ALA D	13.948	49.527	-6.471
	214	414		45 - 71 1	44.754	-4.417	217	778 6	12.788	43.712	-8.875
	217	778 (11.044	43.416	-4.446	237	778 6	12.633	41.111	-4.547
	217	778 1		12.263	41.442	-1.414	217	118 C1	10.473	43.142	-4.870
	217	714 (10.L17	48.293	-4.214	217	448 CO1	10.04	45.993	-3.234
	23.7	111		9.014	45.933	-4.783	217	TTR CES	18.437	47.847	-2.790
	217	779 (8.65-	47.219	-4.381	217	13 817	9.311	47.882	-3.341
	217	778 1		0.753	49.140	-2.911	214	45H W	11.750	41.314	-3.391
40	511	# 3 W S	A	11-645	39.942	-3.227	211	414 C	10.204	21.434	-2.749

				43.347	-1.017	218	05	11.953	>0.340	-8.136
	210	454 0	4.743							
	811	454 E6	14.931	39.544	-2.341	319	41. DD1	34.613	54.704	-1.412
	210	ASH MD2	14.660	39.444	-1.165	210	GLY D	0.478	33.554	-B.209
	211	617 64	8.382	38.131	-2.449	219	SLT C	T. 570	37.304	-3.681
						220	THE W	6.343	24.618	
5	819	6L1 D	7.073	37.60:	-4.876					- 1.203
0	220	THE CL	8.697	35.434	-4.175	220	Pas (4.279	37.044	-0.364
	211	THE	4.417	36.742	-5.911	310	Tat CR	4.825	34.019	-3.926
							7 MB EG2	\$.704	23.694	-1.900
	224	145 BET	4.136	D1.543	-2.451	550				
	217	5 E E D	4.731	30.231	-4.363	537	\$ 2	3.964	39.201	-3.164
	221	344 6	6.760	39.441	-4.303	221	514 0	4.117	40.301	-7.277
	111	11 1 11	3.323	.0.383	-4.544	271	184 05	8.435	46.282	-3.140
	111	mg? m	0.000	31.311	-6.685	272	951 CS	4.471	42.771	-5.173
	111	BET SD	7.748	41.333	-4.993	222	#E7 CG	0.504	41.399	-5.602
10	222	427 66	6.351	40.015	-7.216	222	#ET CA	4.016	39.670	-7.631
										-9.774
	111	MET C	4.877	31.+35	-8.567	\$13	MET 0	7.984	39.967	
	223	ALA W	4.33+	37.244	-8-041	223	ALP CA	6.467	36.010	-1.113
	223	ALA C	5.200	34.044	-0.701	223	ALA D	8.153	35.946	-10.929
						22.	520 6	4.074	34.340	-9.634
	212	ALA ES	6.501	34.907	-7.923					
	114	384 C4	2.758	34.411	-9.702	224	881 6	2.641	37.161	-11.631
	224	111 0	2.141	34.575	-12.557	224	384 68	1.001	B4.995	-8.603
	114	111 06	4.472	34.111	-9.197	225	PRO M	3.154	36.411	-11.111
15							PBC 6			
	225	PRO CA	3.411	34.130	-12.439	225		3.764	34.449	-13.626
	225	*** 0 0	1.404	30.450	-14.804	225	PRD CS	3.653	49.811	-12.854
	115	PRC 66	6.411	40.402	-10.764	221	##3 CD	3.735	34.124	-10.004
					-13.299	224	HIS CA	8.444	36.879	-14.362
	224	MIS W	4.747	37.676						
	224	MIR C	4.418	35.947	-11.041	226	M18 D	4.475	34.809	-14.293
	214	MIS CE		36.046	-13.765	224	MIS CG	7.814	34.859	-13.354
	22.6	M26 WD1	1.040	37.485	-12.170	114	#15 CD2	8.11)	37.118	-14.147
							415 411		37.144	-13.443
20	22.	MIS CES	9.270	30.052	-12.236	114		9.771		
	227	VAL M	3.593	35.366	-14.171	227	APP CO	2.613	34.388	-14.727
	227	TAL C	3.479	35.197	-11.421	127	VAL C	1.016	34.773	-14.490
	117	VAL ES	1.203	37.444	-12.619	127	VAL CEI	1.074	32.474	-14.244
	554	ANT CES	3.204	32.413	-12.871	2 2 3	ALA	1.003	34.242	-14.814
	223	ALA CA	4.011	37.129	-15.517	228	ALA C	0.543	37.534	-16.968
	221	ALA B	-8.213	37.455	-17.828	221	ALA CE	-0.307	34.353	-14.665
	227	SLT N	3.791	31.014	-14.9-1	321	GLY CA	2.352	38.498	-18.231
25	224	GLT C	2.420	37.197	-19.187	229	era o	2.189	37.375	-20.384
23	210	ALA M	2.711	31.711	-16.646	230	ALB EA	2.794	24.001	-19.844
	236	4L4 E	1.424	34.360	-20.133	230	ALE D	1.310	34.263	-21.343
	11:	ALA EB	1.211	31.424	-18.709	231	ALA N	0.263	34.623	-19.324
	231	ALA CA	-L.010	34.414	-19.7-4	231	ALA E	-1.284	31.423	-20.844
	231	ALA D	-1.909	33.814	-21.952	271	ALA CO	-1.932	34.644	-18.549
	132	ALA W	-8.778	34.457	-21.721	232	AL . CA	-1.813	37.663	-21.792
	132	AL . C	-0.201	37.244	-23.078	111	ALA D	-0.841	37.501	-24.187
30	111	ALA CR	-8.742	39.121	-21.377	211	LEU W	0.935	36.724	-22.967
30	233	LBU CA	1.617	34.213	-24.209	233	LEUC	0.021	31.149	-24.886
	4) 1	LEU 0	1.474	35.211	-24.111	2 3 3	LEU CE	3.043	25.877	-23.907
	533	Fån Ce	3.994	34.714	-23.433	\$33	FEG COT	3.219	34.342	-22.921
	233	FEG COS	4.243	37.813	-24.480	534	JL! b	9.337	34.199	-24.047
	23.	ILT CD1	8.304	30.644	-21.437	23.	ILE 161	8,454	31.223	-15.101
	234	1LE CO	-8-811	31.014	-23.570	234	TLE CGE	-1.803	36.900	-24.691
	234	ILP CA	-0.404	32.074	-24.644	234	ILE C	-1.621	33.597	-25.434
35	23.	114 0	-1.413	33.144	-24.344	231	LEU W	-2.390	34.463	-24.77#
30	411	LEL CA	-3.3%	35.021	-25.423	233	LEU C	+3.256	35.843	-26.672
	111						LEU CE			-24.378
		FED 0	-4.109	38. 114	-27.589	133		-4.432	35.765	
	531	LEU CG	-8.140	34.311	-23.34:	233	LEU CO:	-1.652	31.483	-22.149
	235	LEU CD2	-6.212	34.131	-24.120	174	\$13 N	-2.094	34.431	-24.718
	234	680 64	-1.744	37.237	-27.786	114	5 F P C	-1.491	36.392	-29.144
	234	88 D	-1.746	34.634	-30.3at	134	\$ 2 P C P	-0.633	34.234	-27.733
	23.	16. 00	0.571	37.371	-27.982	237	FAR D	-1.044	21.067	-25.682
40	237	LTS CA	-1.144	34.011	-29.952	831	LTS C	-2.113	33.277	-10.248
40	237	LT1 0	-1.374	32.911	-31.444	237	LTS CO	0.272	93.112	-28.553
	237	LTS CG	8.477	33.840	-30.716	237	LT3 CD	1.020	31.935	-30.047

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			2-345			444 454 54	3.525	24.845	-31.594
	111	FAR ES		30.742	-31.774	237 175 02			
	531	#11 E	-2.931	32.989	-20.317	230 #35 64	-4.149	32-141	-21.311
	200	#15 C	-5.314	31.000	-20.447	238 MIS D	-8.713	32.514	-27.562
	111	M16 68	-3.948	30.962	-20.531	134 #11 66	-3.811	29.921	-19.217
								21.111	
	131	mit me:	-1.707	28.679	-21.035	234 WIS CD3	-3.137		-10.394
_	111	m11 CE1	+1.014	20.011	-29.642	230 HZS MF3	-1.948	\$6.418	-00.541
5	211	0 8 D M	-3.641	\$3.927	-21.745	239 PRD Ca	-4.948	34.779	-28.771
	111		-0.204	34.652	-21.572	239 000 0	-1.949	34.519	-27.662
	239	PED CB	-7.018	35.977	-29.713	831 940 CG	-6.646	31.114	-31.027
	234	PED CD	-3.436	331	-30.461	247 68m m	-3.306	32.549	-24.227
	240	ASA CA	-7.521	32.041	-29.216	240 A14 C	-1.500	31.380	-27.960
	2.0	AL- D	-10.340	30.410	-27.576	240 AEN CB	-9.403	31.149	-30.935
						240 414 001	-7.011	31.500	
	840	414 CC	-7.911	30.037	-30.641				-31.147
10	249	454 402	-7.675	87.804	-36.976	\$41 TEP W	-3.354	31.804	-27.244
10	241	TRP CA	-4.304	10.124	-26.126	241 789 C	-9.104	30.431	-24.936
	241	780 0	-1.012	31.933	-24.414	241 707 68	-4.979	29.835	-25.679
		78 P C6				241 TAP COL	-6.330	20.473	
	841		-4.894	20.903	-24.517				
	241	TEP COL	-4.839	28.324	-26.100	241 THP 481	-5.342	27.547	-20.211
	241	TEP CEZ	-4.414	27.474	-27.210	241 TAP CES	-4.097	21.414	-24.981
	241	TRP CII	-3.111	24.784	-27.174	241 789 513	-2.912	27.467	-24.943
	241	TEP ENZ	-3.470		-24.008	242 748 4	-9.717	29.761	-24.142
				16.673					
15	242	THE CA	-10.458	30.119	-22.911	Ses Jan C	-1.447	30.174	-21.747
	242	THE D	-8.333	29.474	-21.937	242 THE EB	-11.570	29.932	-22.675
	242	THE OCI	-10.637	27.786	-22.476	242 THP CG2	-11.494	28.907	-23.811
	243	ASR &	-1.144	39.411	-20.611	243 454 902	-11.717	30.404	-11.747
	143	ASH DD1	-11.465	31.516	-16.768	843 484 CG	-11.043	\$1.171	-17.905
		41= 60							
	243		-9,708	31.130	-11.372	843 ASH CA	-9.853	30.731	-19.044
	843	434 C	-8.487	29.363	-19.010	243 454 D	-7.843	29.134	-11.440
	244	THE L	-7.364	21.362	-19.211	244 THR EA	-9.381	24.734	-19.659
20	244	THE C	-0.133	26.313	-19.862	244 THE D	-7.324	25.757	-19.111
20	244	T## 60	-10.445	24.011	-19.494	244 THE 061	-11.735	24.678	-10.404
	244	THE CG2							
			-10.503	24.515	-19.10		-0.562	24.714	-21.073
	3 - 5	SLM CA	-4.944	26.342	-21.962	343 BLM C	-5.447	27.920	-21.820
	241	BL4 D	-4.572	24.393	-21.447	243 6 LH C8	-7.330	24.5++	-23.297
	2 . 5	810 EG	-0.241	21.526	-23.911	245 GLW CD	-1.473	28.873	-25.420
	241	6L4 DE1	-1.354	24.761	-25.727	245 GL4 MEZ	-7.745	21.312	-14.370
	244	TAL M	-5.697	28.304	-21.211	544 ATT ET	-4.477	29.040	-20.778
25	244	TAL C	-2.934	20.442	-19.467	S.4 AVT D	-2.711	28.227	-19.561
	246	VAL CO	-4.778	10.111	-20.421	244 Val CB3	-3.544	31-272	-28.827
	244	VAL CG2	-3.169	31.131	-21.959	247 ARG 6	-4.761	20.240	-18.442
	247	ARE CA	-4.386	27.714	-17.148	247 ARS C	-1.770	24.212	-17.840
	247	446 D	-2.708		-14.744	247 ARG CO	-1.533	27.447	
				21.005					-14.149
	247	APG CG	-4.987	17.095	-14.012	BAT ARG CD	-4.858	27.179	-13.793
	247	ARE ME	-5.440	26.757	-12.544	247 ARG CZ	-3.813	26.766	-11.815
	247	APE DAS	-7.044	27.484	-11.210	247 485 843	-9.177	24.428	-10.170
30	248	888 4	-4.410	28.505	-10.131	241 SER CA	-4.639	84.131	-10.424
50	24.0	ARR C	-2.457	14.004	-19.075	248 868 0	-1.648	23.253	-10.863
	841	SET CO	-3.034	23.408	-10.372	248 522 05	-4.146	23.010	-14.572
	2++	\$10 m	-2.300	24.883	-20.136	349 BBR CA	-1.713	24.874	-20.053
	249	814 (-0.071	25.302	-19.942	247 528 8	1. U.	24.705	-20.049
	145	312 63	-1.369	28.758	-22.040	249 520 06	- 5. 300	25.419	-22.954
	210	LEUR	-1.200		-19.140	230 LOU CO2	1.824	29.914	
				26.333					-11.122
	110	TEN COT	-0.373	30.453	-17.200	290 F20 CC	4.312	21.431	-14.511
35	210	TEN CB	8.178	20.041	-17.863	\$30 FER CV	0.718	24.637	-18.214
	210	LEU C	1.092	28.494	-17.265	890 LEU C	1.203	25.421	-17.032
	251	5L % &	0.044	20.027	-14.714	251 BLW ME2	-2.710	25.012	-12.337
	211	6L4 DE2	-2.019	23.424	-12.738	291 BLN CD	-2.345	24.850	-13.634
	131	BLW CG	-1.216	24.414	-11.114		-9.987	13.421	-14.877
	251	GLW CA	0.301	23.941	-10.745	891 BLM C	4.939	22.044	-16.361
	401	SLH D	1.743	22.01-	-15.414	282 AS4 M	0.433	51.304	-17.800
	111	454 64	1.002	21.704	-18.282	202 ASH C	2.194	21.359	-10.001
40	111	ASN C	2.101	20.442	-11.760	BIZ ASH CO	0.004	26.780	-19.212
-	111	45 H & 6	-1.034		-19.573	882 48% 8D1		19.195	
	472	-3	-1.44	19.924	-38.213	ere =>= 851	-4.834	44.22	-17.582

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					-17.161	253	T 88 6	3.614	22.805	-18.921
	2 5 2	434 MG2	-2.234	29.874	-19.713	155	THE E	9.301	23.247	-18.811
	213	101 64	4.234	25.733	-19.421	211	43 ENT	4.911	23.672	-21.952
	213	7-1 0	6.741	24.937	-10.422	153	141 662	3.147	23.130	-32.032
	213	fa1 861	3.111			234	THE CA	4.214	23.412	-14.558
	864	1=4 4	8.211	23.377	-17.151	234	Te4 D	7.402	21.550	-17.091
5	254	4 m h C	7.444	22.720	-14.413	234	Tet 053	9.121	22,170	-16.040
J	21-	THE CO	5.664	23.998	-13.131		THE SES	3.411	23.294	-14.874
	254	1m3 C62	4.530	24.549	-10.807	255	THE	9.021	22.031	-14.414
	211	1=1 C1	9.771	22.19.	-11.817	255		11.065	23.411	-15.697
	234	THE O	9.414	22.746	-13.474	255	T=1 [1		22.619	-13.006
	311	TAR 061	21.002	23.707	-17.321	253	4=> CES	12.214		
	214	L42 P	9.601	80.702	-14.314	256	LTS CA	9.144	20.043	-13.616
	254	LTEC	10.31:	20.333	-12.063	254	LTS D	11.662	20.274	-11.492
10	214	LTS CT	9.02+	10.540	-11.249	234	LYS CC	9.016	17.805	-31.971
10	254	LTS CO	10.214	16.948	-11.771	254	LTJ ER	10.217	39.940	-10.621
	254	1.72 M2	0.243	14.869	-11.034	257	FED #	10.212	80.474	-10.674
	211	LEU CA	11.272	21.036	-9.513	257	Ltu C	11.250	20.232	-8.614
	217	LIL D	12.094	20.843	-7.737	257	LEU ER	11.107	22.947	-4.811
	257	LEU CC	11.357	23.470	-10.844	211	Ltu CD1	11.243	25.003	-9.921
	217	LEU CCI	32.678	23.441	-11.325	251	SLT #	10.631	39.232	-8.271
	111	BLT CA	16.665	11.713	-4.879	254	SLT C	9.148	18.703	-4.373
	254	614 0	6.213	18.954	-7.252	231	45 P M	9.824	10.202	-5.150
15	211	457 EA	7.737	17.494	-4.514	299	ASP C	4.411	28.941	-4.789
	231	450 D	4.453	10.0>+	-4.234	211	ASP CB	7.414	17.940	-3.611
	231	450 EE	4.701	17.126	-2.243	251	41+ DD1	\$.611	17.527	-2.154
	217	45" 002	7.011	14.297	-1.321	240	5 E+ h	4.540	14.610	-5.312
	Pa t	111 61	6.481	19.507	-1.525	240	111 C	4.044	20.342	-0.211
	245	\$ t + D	3.586	21.513	-4,444	240	111 61	3.345	18.919	-4.215
	265	310 00	2.743	17.937	-5.4+1	241	PHE N	4.241	11.778	-3.112
		9H(CA	3.831	21.461	-1.065	261	PHE C	4.544	21.846	-1.063
20	8 1		3.744	21.041	-1.432	341	PHE CS	4.057	19.769	-0.943
	161	741 B	1.547	20.337	0.719	261	P## CD3	2.204	20.163	1.111
	263	Pet CG		21.860	1.555	241	PHE CEL	1.717	28.717	2.315
	241	but cos	3.945	21.602	2.748	241	PHE CI	2.405	21.461	3.114
	261	7#1 CE2	8.774	21.793	-2.303	2 4 2	118 64	6.613	22.914	-2.251
	2 6 2		6.826	23.619	-3.945	167	TYR D	7.701	24.953	-3.213
	247		4.122	22.433	-1.031	367	778 CG	8.104	21.892	-8.454
	341	TYR C8	9.054	20.434	-0.34	207	TYR CD2	6.141	22.641	0.451
25	342	TTR CC3	8.842	19.073	4.432	202	TTR CEZ	8.114	22.049	1.942
	247	778 663	8.047	20.671	2.018	262	118 D=	7.945	20.029	3.201
	242	110 (1	4.624	23.104	-4.473	243	794 C.	6.112	11.451	-4.822
	363	TTR 4		23.610	-4.954	265	TVE S	8.783	24.117	-8.111
	243	778 (8.624	22.765	-4.481	243	TYR CC	9.219	23.035	-4.648
	243	170 CB	7.121		-4.557	243	110 502	9.800	21.342	-4.993
	243	T## 603	10.064	24.046	-4.141	245	110 682	33.042	21.440	-4.491
	243	778 681			-1.154	201	TTR D	17.045	23.949	-4.817
30	243	718 63	\$1.636	23.418	-4.516	111	817 64	3.301	21.044	-7.412
	244	61.	4.471	23.141	-1.114	111	614 0	4.647	21.274	-1.343
	244	641 6	3.647	22-194	-1.15	111	LTS CA	3.634	21.711	-10.971
	261	175 4	3.431	82.477	-11.444	145	L71 D	5.414	21.943	-12.384
	241	L75 C	9.388	21.232	-12.044	265	LTS EE	1.491	21.963	-11.308
	245	LTS CB	2.755	\$2.671		311	LTS CE	-9.612	10.494	-11.391
	449	LTS CD	0.710	30.541	-12.679	111	617 8	8.787	23.224	-10.917
	265	L 7 6 42	-1.478	23.797	-11.323	761	617 6	7.133	25.012	-11.010
35	264	BLT CA	7.120	83.412		267	LEU W	B. 242	11.33.	-12.490
	400	SLT D	4-377	81.793	-11.648	267		7.904	26.771	-14.417
	267	FEO CE	8.491	24.440	-11.097	267	LTV C	10.010	24.933	-13.214
	247	LEU D	7.913	23.909	-13.298		110 (8	10.074	27.373	-11.215
	247	LIU CG	10.412	11.515	-14.018	267	140 601	7.044	27.863	-10.632
	247	FIN CDS	11.924	37.923	-14.327	141	ILT N	7.424	30.244	-17.005
	341	ILE CA	4.404	28.033	-15.944	241	114 CD	8.767	20.210	-11.811
	201	111 0	8.519	88.713	-14.912	201	114 (62	4.14)	20.925	-14.967
40	8 + 1	14 661	6.091	30.541	-15.543	1.1		7.107	27.1.3	-11.217
	3 4 5	ILE CD1	8.391	31.745	-14.262		654 h	****		

	249	884 64	808.7	27.475	-11.437	211	41. 5	.e. 89 b	28.954	-018-695
	444	46- 0	1.045	27.562	-76.44.2	244	AIN CE	b. 4 7 /	14.613	
	20.7	41: 66	1.101		-21.215			0.113		-: 4,431
				20.451		241	9 8 4 83 T		17.414	2.12:
	247	8801	11.011	25.706	-11.072	272	AVE P	4 908	116.01	-26.786
	170	WAL GA	3.313		-21-616	210	WAL A	A. 851	\$8.997	- 1.434
	270	WAL D	8.017	27.969	-23.572	317	VAL CE	3.646	F1.710	-21-627
5 .	876	TAL EGS	6.847	32.717	-21.676	275	VAL CER	3.6/0	F1.362	. 13.73.
-	171		9.323	29.751	-23.331	111	61.4 64	7.857	29.210	
										-14,541
	273	\$ L 3 :	6.469	2/- 71 +	-21-531	811	EL . D	4.213	27.864	- 14 . 0 9 .
	17:	SLa C1	4.104	34.120	-34-944	8"1	BL4 EG	0.484	£4.414	-14.785
	811	SLW ED	\$0 . Bb \$	28.865	-11.982	271	er# 0[]	11.344	18.574	-17.716
	271	614 412	15.802	26.513	-21 -116	272	ALS N	3.077	24.000	-74.897
	272	ALA EA	6.8.4	2 7 . 2	-24.445	2"2	ALA E	771	87.384	-14.76
	172	ALA D	3.114	23.501	-21.16:	272	ALA EP	6.743	24. 742	- 12 .). \$ 2
10	11)	81.0 13	4.1.7		- 25 - 3 3:			4.1.0		
				24.461		2 - 3	ALP EA		- Be-383	-12.55
	113	AL . C	4 - 847	27.628	-2620	£^3	8L4 D	7.348	47.219	#44.5°\$5
	213	ALA CS	4.136	27.773	-1111	2 :-	ale m	1.785	48.464	- 14.74/
	274	ALG CB	3.452	33.391	-26.218	274	ALS CO	2.189	29.166	-48. FAT
	274	4L4 C	2.730	21.347	-27.095	2 74	ALA 3	9.969	20.749	
	275	614 %	3.110	27.194	-2' -314	2.3	614 Se	4.948	24.309	- 18 227
	171	SLW C	1.1.1	27.261	17.777	155	614 0	1.740	211.067	
					- ·					- 7.516
15	273	614 01	3 . 1 3 1	21.332	• 3 0 • 0	875	SL4 CB	4	28.776	10 . 5 2 0
, ,	275	SLA CL	9 - 8 3 1	23.644	- 27 . 4 47	7"+	ero el	-3.473	23.434	11 . 63 ≥
	273	GLA DII	-1.174	23.143	. 24.729	2 3	ALK MIZ	-1.1:3	.73.411	-16.536

The above structural studies together with the kinetic data presented herein and elsewhere (Philipp, M., et al. (1983) Mol. Cell. Biochem. 51, 5-32; Svendsen, I.B. (1976) Carlsberg Res. Comm. 41, 237-291; Markland, S.F. Id; Stauffe, D.C., et al. (1965) J. Biol. Chem. 244, 5333-5338) indicate that the subsites in the binding cleft of subtilisin are capable of interacting with substrate amino acid residues from P-4 to P-2'.

The most extensively studied of the above residues are Gly166, Gly169 and Ala152. These amino acids were identified as residues within the S-1 subsite. As seen in Fig. 3, which is a stereoview of the S-1 subsite, Gly166 and Gly169 occupy positions at the bottom of the S-1 subsite, whereas Ala152 occupies a position near the top of S-1, close to the catalytic Ser221.

All 19 amino acid substitutions of Gly166 and Gly169 have been made. As will be indicated in the examples which follow, the preferred replacement amino acids for Gly166 and/or Gly169 will depend on the specific amino acid occupying the P-1 position of a given substrate.

The only substitutions of Ala152 presently made and analyzed comprise the replacement of Ala152 with Gly and Ser. The results of these substitutions on P-1 specificity will be presented in the examples.

In addition to those residues specifically associated with specificity for the P-1 substrate amino acid, Tyr104 has been identified as being involved with P-4 specificity. Substitutions at Phe189 and Tyr217, however, are expected to respectively effect P-2' and P-1' specificity.

The catalytic activity of subtilisin has also been modified by single amino acid substitutions at Asn155. The catalytic triad of subtilisin is shown in Fig. 4. As can be seen, Ser221, His64 and Asp32 arc positioned to facilitate nucleophilic attach by the serine hydoxylate on the carbonyl of the scissile peptide bond. Crystallographic studies of subtilisin (Robertus, et al. (1972) Biochem. 11, 4293-4303; Matthews, et al. (1975) J. Biol. Chem. 250, 7120-7126; Poulos, et al. (1976) J. Biol. Chem. 250, 1097-1103) show that two hydrogen bonds are formed with the oxyanion of the substrate transition state. One hydrogen bond donor is from the catalytic serine-221 main-chain amide while the other is from one of the NE2 protons of the asparagine-155 side chain. See Fig. 4.

Asn155 was substituted with Ala, Asp, His, Glu and Thr. These substitutions were made to investigate the the stabilization of the charged tetrahedral intermediate of the transition state complex by the potential hydrogen bond between the side chain of Asn155 and the oxyanion of the intermediate. These particular substitutions caused large decreases in substrate turnover, kcat (200 to 4,000 fold), marginal decreases in substrate binding Km (up to 7 fold), and a loss in transition state stabilization energy of 2.2 to 4.7 kcal/mol. The retention of Km and the drop in kcat will make these mutant enzymes useful as binding proteins for specific; peptide sequences, the nature of which will be determined by the specificity of the precursor protease.

Various other amino acid residues have been identified which affect alkaline stability. In some cases, mutants having altered alkaline stability also have altered thermal stability.

In <u>B</u> amyloliquefaciens subtilisin residues Asp36, Ile107, Lys170, Ser204 and Lys213 have been identified as residues which upon substitution with a different amino acid alter the alkaline stability of the mutated enzyme as compared to the precursor enzyme. The substitution of Asp36 with Ala and the substitution of Lys170 with Glu each resulted in a mutant enzyme having a lower alkaline stability as compared to the wild type subtilisin. When Ile107 was substituted with Val, Ser204 substituted with Cys, Arg or Leu or Lys213 substituted with Arg, the mutant subtilisin had a greater alkaline stability as compared

to the wild type subtilisin. However, the mutant Ser204P demonstrated a decrease in alkaline stability.

In addition, other residues, identified as being associated with the modification of other properties of subtilisin, also affect alkaline stability. These residues include Ser24, Met50, Glu156, Gly166, Gly169 and Tyr217. Specifically the following particular substitutions result in an increased alkaline stability: Ser24C, Met50F, Gly156Q or S, Gly166A, H, K, N or Q, Gly169S or A, and Tyr217F, K, R or L. The mutant Met50V, on the other hand, results in a decrease in the alkaline stability of the mutant subtilisin as compared to wild type subtilisin.

Other residues involved in alkaline stability based on the alkaline stability screen include Asp197 and Met222. Particular mutants include Asp197(R or A) and Met 222 (all other amino acids).

Various other residues have been identified as being involved in thermal stability as determined by the thermal stability screen herein. These residues include the above identified residues which effect alkaline stability and Met199 and Tyr21. These latter two residues are also believed to be important for alkaline stability. Mutants at these residues include I199 and F21.

The amino acid sequence of <u>B</u>. <u>amyloliquefaciens</u> substilisin has also been modified by substituting two or more amino acids of the wild-type sequence. Six categories of multiply substituted mutant subtilisin have been identified. The first two categories comprise thermally and oxidatively stable mutants. The next three other categories comprise mutants which combine the useful properties of any of several single mutations of <u>B</u>. <u>amyloliquefaciens</u> subtilisin. The last category comprises mutants which have modified alkaline and/or thermal stability.

The first category comprises double mutants in which two cysteine residues have been substituted at various amino acid residue positions within the subtilisin molecule. Formation of disulfide bridges between the two substituted cysteine residues results in mutant subtilisins with altered thermal stability and catalytic activity. These mutants include A21/C22/C87 and C24/C87 which will be described in more detail in Example 11.

The second category of multiple subtilisin mutants comprises mutants which are stable in the presence of various oxidizing agents such as hydrogen peroxide or peracids. Examples 1 and 2 describe these mutants which include F50/I124/Q222, F50/I124, F50/Q222, F50/L124/Q222, I124/Q222 and L124/Q222.

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The third category of multiple subtilisin mutants comprises mutants with substitutions at position 222 combined with various substitutions at positions 166 or 169. These mutants, for example, combine the property of oxidative stability of the A222 mutation with the altered substrate specificity of the various 166 or 169 substitutions. Such multiple mutants include A166/A222, A166/C222, F166/C222, K166/A222, K166/C222, V166/A222 and V166/C222. The K166/A222 mutant subtilisin, for example, has a kcat/Km ratio which is approximately two times greater than that of the single A222 mutant subtilisin when compared using a substrate with phenylalanine as the P-1 amino acid. This category of multiple mutant is described in more detail in Example 12.

The fourth category of multiple mutants combines substitutions at position 156 (Glu to Q or S) with the substitution of Lys at position 166. Either of these single mutations improve enzyme performance upon substrates with glutamate as the P-1 amino acid. When these single mutations are combined, the resulting multiple enzyme mutants perform better than either precursor. See Example 9.

The fifth category of multiple mutants contain the substitution of up to four amino acids of the B. <u>amyloliquefaciens</u> subtilisin sequence. These mutants have specific properties which are virtually identicle to the properties of the subtilisin from B. <u>licheniformis</u>. The subtilisin from B. <u>licheniformis</u> differs from B. <u>amyloliquefaciens</u> subtilisin at 87 out of 275 amino acids. The multiple mutant F50/S156/A169/L217 was found to have similar substrate specificity and kinetics to the licheniformis enzyme. (See Example 13.) However, this is probably due to only three of the mutations (S156, A169 and L217) which are present in the substrate binding region of the enzyme. It is quite surprising that, by making only three changes out of the 87 different amino acids between the sequence of the two enzymes, the B. <u>amyloliquifaciens</u> enzyme was converted into an enzyme with properties similar to B. <u>licheniformis</u> enzyme. Other enzymes in this series include F50/Q156/N166/L217 and F50/S156/L217.

The sixth category of multiple mutants includes the combination of substitutions at position 107 (lle to V) with the substitution of Lys at position 213 with Arg, and the combination of substitutions of position 204 (preferably Ser to C or L but also to all other amino acids) with the substitution of Lys at position 213 with R. Other multiple mutants which have altered alkaline stability include Q156/K166, Q156/N166, S156/K166, S156/K166, S156/N166 (previously identified as having altered substrate specificity), and F50/S156/A169/L217 (previously identified as a mutant of B. amyloliquifaciens subtilisin having properties similar to subtilisin from B. licheniformis). The mutant F50/V107/R213 was constructed based on the observed increase in alkaline stability for the single mutants F50, V107 and R213. It was determined that the V107/R213 mutant had an increased alkaline stability as compared to the wild type subtilisin. In this particular mutant, the increased

alkaline stability was the result of the cumulative stability of each of the individual mutations. Similarly, the mutant F50/V107/R213 had an even greater alkaline stability as compared to the V107/R213 mutant indicating that the increase in the alkaline stability due to the F50 mutation was also cumulative.

Table IV summarizes the multiple mutants which have been made including those not mentioned above. In addition, based in part on the above results, substitution at the following residues in subtilisin is expected to produce a multiple mutant having increased thermal and alkaline stability: Ser24, Met50, Ile107, Glu156, Gly169, Gly169, Ser204, Lys213, Gly215, and Tyr217.

TABLE IV

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Double Mutants	Triple, Quadruple or Other Multiple
C22/C87	F50/I124/Q222
C24/C87	F50/L124/Q222
V45/V48	F50/L124/A222
C49/C94	A21/C22/C87
C49/C95	F50/S156/N166/L217
C50/C95	F50/Q156/N166/L217
C50/C110	F50/S156/A169/L217
F50/l124	F50/S156/L217
F50/Q222	F50/Q156/K166/L217
I124/Q222	F50/S156/K166/L217
Q156/D166	F50/Q156/K166/K217
Q156/K166	F50/S156/K166/K217
Q156/N166	F50/V107/R213
S156/D166	[S153/S156/A158/G159/S160/Δ161-164/I165/S166/A169/R170]
S156/K166	
S156/N166	L204/R213
S156/A169	R213/204A, E, Q, D, N, G, K, V, R, T, P, I, M, F, Y, W or H
A166/A222	
A166/C222	
F166/A222	V107/R213
F166/C222	
K166/A222	
K166/C222	
V166/A222	
V166/C222	
A169/A222	
A169/A222	
A169/C222	
A21/C22	

In addition to the above identified amino acid residues, other amino acid residues of subtilisin are also considered to be important with regard to substrate specificity. Mutation of each of these residues is expected to produce changes in the substrate specificity of subtilisin. Moreover, multiple mutations among these residues and among the previously identified residues are also expected to produce subtilisin mutants having novel substrate specificity.

Particularly important residues are His67, Ile107, Leu126 and Leu135. Mutation of His67 should alter the S-1' subsite, thereby altering the specificity of the mutant for the P-1' substrate residue. Changes at this position could also affect the pH activity profile of the mutant. This residue was identified based on the inventor's substrate modeling from product inhibitor complexes.

lle107 is involved in P-4 binding. Mutation at this position thus should alter specificity for the P-4 substrate residue in addition to the observed effect on alkaline stability. Ile107 was also identified by molecular modeling from product inhibitor complexes.

The S-2 binding site includes the Leu126 residue. Modification at this position should therefore affect P-2 specificity. Moreover, this residue is believed to be important to convert subtilisin to an amino peptidase.

The pH activity profile should also be modified by appropriate substitution. These residues were identified from inspection of the refined model, the three dimensional structure from modeling studies. A longer side chain is expected to preclude binding of any side chain at the S-2 subsite. Therefore, binding would be restricted to subsites S-1, S-1', S-2', S-3' and cleavage would be forced to occur after the amino terminal peptide.

Leu135 is in the S-4 subsite and if mutated should alter substrate specificity for P-4 if mutated. This residue was identified by inspection of the three-dimensional structure and modeling based on the product inhibitor complex of F222.

In addition to theses sites, specific amino acid residues within the segments 97-103, 126-129 and 213-215 are also believed to be important to substrate binding.

Segments 97-103 and 126-129 form an antiparallel beta sheet with the main chain of substrate residues P-4 through P-2. Mutating residues in those regions should affect the substrate orientation through main chain (enzyme) - main chain (substrate) interactions, since the main chain of these substrate residues do not interact with these particular residues within the S-4 through S-2 subsites.

Within the segment 97-103, Gly97 and Asp99 may be mutated to alter the position of residues 101-103 within the segment. Changes at these sites must be compatible, however. In <u>B. amyloliquifaciens</u> subtilisin Asp99 stabilizes a turn in the main chain tertiary folding that affects the direction of residues 101-103. <u>B. licheniformis subtilisin Asp97, functions in an analogous manner.</u>

In addition to Gly97 and Asp99, Ser101 interacts with Asp99 in B. amyliquefaciens subtilisin to stabilize the same main chain turn. Alterations at this residue should after the 101-103 main chain direction. Mutations at Glu103 are also expected to affect the 101-103 main chain direction.

The side chain of Gly102 interacts with the substrate P-3 amino acid. Side chains of substituted amino acids thus are expected to significantly affect specificity for the P-3 substrate amino acids.

All the amino acids within the 127-129 segment are considered important to substrate specificity. Gly127 is positioned such that its side chain interacts with the S-1 and S-3 subsites. Altering this residue thus should alter the specificity for P-1 and P-3 residues of the substrate.

The side chain of Gly128 comprises a part of both the S-2 and S-4 subsites. Altered specificity for P-2 and P-4 therefore would be expected upon mutation. Moreover, such mutation may convert subtilisin into an amino peptidase for the same reasons substitutions of Leu126 would be expected to produce that result.

The Pro129 residue is likely to restrict the conformational freedom of the sequence 126-133, residues which may play a major role in determining P-1 specificity. Replacing Pro may introduce more flexibility thereby broadening the range of binding capabilities of such mutants.

The side chain of Lys213 is located within the S-3 subsite. All of the amino acids within the 213-215 segment are also considered to be important to substrate specificity. Accordingly, altered P-3 substrate specificity is expected upon mutation of this residue.

The Tyr214 residue does not interact with substrate but is positioned such that it could affect the conformation of the hair pin loop 204-217.

Finally, mutation of the Gly215 residue should affect the S-3' subsite, and thereby alter P-3' specificity.

In addition to the above substitutions of amino acids, the insertion or deletion of one or more amino acids within the external loop comprising residues 152-172 may also affect specificity. This is because these residues may play a role in the "secondary contact region" described in the model of streptomyces subtilisin inhibitor complexed with subtilisin. Hirono, et al. (1984) J. Mol. Biol. 178, 389-413. Thermitase K has a deletion in this region, which eliminates several of these "secondary contact" residues. In particular, deletion of residues 161 through 164 is expected to produce a mutant subtilisin having modified substrate specificity. In addition, a rearrangement in this area induced by the deletion should after the position of many residues involved in substrate binding, predominantly at P-1. This, in turn, should affect overall activity against proteinaceous substrates

The effect of deletion of residues 161 through 164 has been shown by comparing the activity of the wild type (WT) enzyme with a mutant enzyme containing this deletion as well as multiple substitutions (i.e., \$153/\$158/A158/G159/\$160/\Delta164/165/\$166/A169/R170). This produced the following results:

TABLE V

	kcat	Km	kcat/Km
WT	50	1.4x10 ⁻⁴	3.6x10 ⁵
Deletion mutant	8	5.0x10 ⁻⁶	1.6x10 ⁶

The WT has a kcat 3 times greater than the deletion mutant but substrate binding is 28 fold tighter by the deletion mutant. The overall efficiency of the deletion mutant is thus 4.4 times higher than the WT enzyme.

All of these above identified residues which have yet to be substituted, deleted or inserted into are presented in Table VI.

TABLE VI

	Substitution/In	sertion/Deletion
	Res	idues
	His67	Ala152
	Leu126	Ala153
1	Leu135	Gly154
	Gly97	Asn155
ı	Asp99	Gly156
	Ser101	Gly157
	Gly102	Gly160
	Glu103	Thr158
	Leu126	Ser159
	Gly127	Ser161
	Gly128	Ser162
ı	Pro129	Ser163
	Tyr214	Thr164
	Gly215	Val165
	Gly166	Gly169
	Tyr167	Lys170
	Pro168	Tyr171
		Pro172

The following disclosure is intended to serve as a representation of embodiments herein, and should not be construed as limiting the scope of this application. These specific examples disclose the construction of certain of the above identified mutants. The construction of the other mutants, however, is apparent from the disclosure herein and that presented in EPO Publication No. 0130756.

All literature citations are expressly incorporated by reference.

EXAMPLE 1

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Identification of Peracid Oxidizable Residues of Subtilisin Q222 and L222

As shown in Figures 6A and 6B, organic peracid oxidants inactivate the mutant subtilisins Met222L and Met222Q (L222 and Q222). This example describes the identification of peracid oxidizable sites in these mutant subtilisins.

First, the type of amino acid involved in peracid oxidation was determined. Except under drastic conditions (Means, G.E., et al. (1971) Chemical Modifications of Proteins, Holden-Day, S.F., CA, pp. 160-162), organic peracids modify only methionine and tryptophan in subtilisin. Difference spectra of the enzyme over the 250nm to 350nm range were determined during an inactivation titration employing the reagent, diperdodecanoic acid (DPDA) as oxidant. Despite quantitative inactivation of the enzyme, no change in absorbance over this wavelength range was noted as shown in Figures 7A and 7B indicating that tryptophan was not oxidized. Fontana, A., et al. (1980) Methods in Peptide and Protein Sequence Analysis - (C. Birr ed.) Elsevier, New York, p. 309. The absence of tryptophan modification implied oxidation of one or more of the remaining methionines of B. amyloliquefaciens subtilisin. See Figure 1.

To confirm this result the recombinant subtilisin Met222F was cleaved with cyanogen bromide (CNBr) both before and after oxidation by DPDA. The peptides produced by CNBr cleavage were analyzed on high resolution SDS-pyridine peptide gels (SPG).

Subtilisin Met222F (F222) was oxidized in the following manner. Purified F222 was resuspended in 0.1 M sodium borate pH 9.5 at 10 mg/ml and was added to a final concentration of 26 diperdodecanoic acid

(DPDA) at 26 mg/ml was added to produce an effective active oxygen concentration of 30 ppm. The sample was incubated for at least 30 minutes at room temperature and then quenched with 0.1 volume of 1 M Tris pH 8.6 buffer to produce a final concentration of 0.1 M Tris pH 8.6). 3mM phenylmethylsulfonyl fluoride (PMSF) was added and 2.5 ml of the sample was applied to a Pharmacia PD10 column equilibrated in 10 mM sodium phosphate pH 6.2, 1 mM PMSF. 3.5 ml of 10 mM sodium phosphate pH6.2, 1mM PMSF was applied and the eluant collected.

F222 and DPDA oxidized F222 were precipitated with 9 volumes of acetone at -20 °C. The samples were resuspended at 10 mg/ml in 8M urea in 88% formic acid and allowed to sit for 5 minutes. An equal volume of 200 mg/ml CNBr in 88% formic acid was added (5 mg/ml protein) and the samples incubated for 2 hours at room temperature in the dark. Prior to gel electrophoresis, the samples were lyophilized and resuspended at 2-5 mg/ml in sample buffer (1% pyridine, 5% NaDodSO₄, 5% glycerol and bromophenol blue) and disassociated at 95 °C for 3 minutes.

The samples were electrophoresed on discontinuous polyacrylamide gels (Kyte, J., et al. (1953) Anal. Bioch. 133, 515-522). The gels were stained using the Pharmacia silver staining technique (Sammons, D.W., et al. (1981) Electrophoresis 2 135-141).

The results of this experiment are shown in Figure 8. As can be seen, F222 treated with CNBr only gives nine resolved bands on SPG. However, when F222 is also treated with DPDA prior to cleavage, bands X, 7 and 9 disappear whereas bands 5 and 6 are greatly increased in intensity.

In order to determine which of the methionines were effected, each of the CNBr peptides was isolated by reversed phase HPLC and further characterized. The buffer system in both Solvent A (aqueous) and Solvent B (organic) for all HPLC separations was 0.05% triethylamime/trifloroacetic acid (TEA-TFA). In all cases unless noted, solvent A consisted of 0.05% TEA-TFA in H₂0, solvent B was 0.05% TEA-TFA in 1-propanol, and the flow rate was 0.5 ml/minute.

For HPLC analysis, two injections of 1 mg enzyme digest were used. Three samples were acetone precipitated, washed and dried. The dried 1 mg samples were resuspended at 10 mg/ml in 8M urea, 88% formic acid; an equal volume of 200 mg/ml CNBr in 88% formic acid was added (5 mg/ml protein). After incubation for 2 hours in the dark at room temperature, the samples were desalted on a 0.8 cm X 7 cm column of Tris Acryl GF05 coarse resin (IBF, Paris, France) equilibrated with 40% solvent B, 60% solvent A. 200 ul samples were applied at a flow rate of 1 ml a minute and 1.0-1.2 ml collected by monitoring the absorbance at 280nm. Prior to injection on the HPLC, each desalted sample was diluted with 3 volumes of solvent A. The samples were injected at 1.0 ml/min (2 minutes) and the flow then adjusted to 0.5 ml/min (100% A). After 2 minutes, a linear gradient to 60% B at 1.0% B/min was initiated. From each 1 mg run, the pooled peaks were sampled (50ul) and analyzed by gel electrophoresis as described above.

Each polypeptide isolated by reversed phase HPLC was further analyzed for homogeneity by SPG. The position of each peptide on the known gene sequence (Wells, J.A., et al. (1983) <u>Nucleic Acids Res. 11</u> 7911-7924) was obtained through a combination of amino acid compositional analysis and, where needed, amino terminal sequencing.

Prior to such analysis the following peptides were to rechromatographed.

40 1. CNBr peptides from F222 not treated with DPDA:

Peptide 5 was subjected to two additional reversed phase separations. The 10 cm C4 column was equilibrated to 80%A/ 20%B and the pooled sample applied and washed for 2 minutes. Next an 0.5% ml B/min gradient was initiated. Fractions from this separation were again rerun, this time on the 25 cm C4 column, and employing 0.05% TEA-TFA in acetonitrile/1-propanol (1:1) for solvent B. The gradient was identical to the one just described.

Peptide "X" was subjected to one additional separation after the initial chromatography. The sample was applied and washed for 2 minutes at 0.5ml/min (100%A), and a 0.5% ml B/min gradient was initiated.

Peptides 7 and 9 were rechromatographed in a similar manner to the first rerun of peptide 5.

Peptide 8 was purified to homogeneity after the initial separation.

2. CNBr Peptides from DPDA Oxidized F222:

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Peptides 5 and 6 from a CNBr digest of the oxidized F222 were purified in the same manner as peptide 5 from the untreated enzyme.

Amino acid compositional analysis was obtained as follows. Samples (-1nM each amino acid) were dried, hydrolyzed in vacuo with 100 ul 6N HCl at 106 °C for 24 hours and then dried in a Speed Vac. The samples were analyzed on a Beckmann 6300 AA analyzer employing ninhydrin detection.

Amino terminal sequence data was obtained as previously described (Rodriguez, H., et al. (1984) Anal. Biochem. 134, 538-547).

The results are shown in Table VII and Figure 9.

TABLE VII

Amino and CC	OOH terminii of CNBr fragm	ents Terminus and Method
Fragment	amino, method	COOH, method
×	1, sequence	50, composition
9	51, sequence	119, composition
7	125, sequence	199, composition
8	200, sequence	275, composition
5ox	1, sequence	119, composition
6ox	120, composition	199, composition

Peptides 5ox and 6ox refer to peptides 5 and 6 isolated from CNBr digests of the oxidized protein where their respective levels are enhanced.

From the data in Table VII and the comparison of SPG tracks for the oxidized and native protein digests in Figure 8, it is apparent that (1) Met50 is oxidized leading to the loss of peptides X and 9 and the appearance of 5; and (2) Met124 is also oxidized leading to the loss of peptide 7 and the accumulation of peptide 6. Thus oxidation of B. amyloliquifaciens subtilisin with the peracid, diperdocecanoic acid leads to the specific oxidation of methionine at residues 50 and 124.

EXAMPLE 2

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Substitution at Met50 and Met124 in Subtilisin Met222Q

The choice of amino acid for substitution at Met50 was based on the available sequence data for subtilisins from B. licheniformis (Smith, E.C., et al. (1968) J. Biol. Chem. 243, 2184-2191), B.DY (Nedkov, P., et al. (1983) Hoppe Sayler's Z. Physiol. Chem. 364 1537-1540), B. amylosacchariticus (Markland, F.S., et al. (1967) J. Biol. Chem. 242 5198-5211) and B. subtilis (Stahl, M.L., et al. (1984) J. Bacteriol. 158, 411-418). In all cases, position 50 is a phenylalanine. See Figure 5. Therefore, Phe50 was chosen for construction.

At position 124, all known subtilisins possess a methionine. See Figure 5. Molecular modelling of the x-ray derived protein structure was therefore rehired to determine the most probable candidates for substitution. From all 19 candidates, isoleucine and leucine were chosen as the best residues to employ. In order to test whether or not modification at one site but not both was sufficient to increase oxidative stability, all possible combinations were built on the Q222 backbone (F50/Q222, I124/Q222, F50/I124/Q222).

A. Construction of Mutations Between Codons 45 and 50

All manipulations for cassette mutagenesis were carried out on pS4.5 using methods disclosed in EPO Publication No. 0130756 and Wells, J.A., et al, (1985) Gene 34, 315-323. The p∆50 in Fig. 10, line 4, mutations was produced using the mutagenesis primer shown in Fig. 10, line 6, and employed an approach designated as restriction-purification which is described below. Briefly, a M13 template containing the subtilisin gene, M13mp11-SUBT was used for heteroduplex synthesis (Adelman, et al (1983), DNA 2, 183-193). Following transfection of JM101 (ATCC 33876), the 1.5 kb EcoRI-BamHI fragment containing the subtilisin gene was subcloned from M13mp11 SUBT rf into a recipient vector fragment of pBS42 the construction of which is described in EPO Publication No. 0130756. To enrich for the mutant sequence (p∆50, line 4), the resulting plasmid pool was digested with Kpnl, and linear molecules were purified by polyacrylamide gel electrophoresis. Linear molecules were ligated back to a circular form, and transformed into E. coli MM294 cells (ATCC 31446). Isolated plasmids were screened by restriction analysis for the KpnI, site. KpnI⁺ plasmids were sequenced and confirmed the p∆50 sequence. Asterisks in Figure 11 indicate the bases that are mutated from the wid type sequence (line 4). p 50 (line 4) was cut with Stul and EcoRI and the 0.5 Kb tragment containing the 5' half of the subtilisin gene was purified (fragment 1). pΔ50 (line 4) was digested with KonI and EcoRI and the 4.0 Kb fragment containing the 3' half of the subtilisin gene and vector sequences was purified (fragment 2). Fragments 1 and 2 (line 5), and duplex DNA

cassettes coding for mutations desired (shaded sequence, line 6) were mixed in a molar ratio of 1:1:10, respectively. For the particular construction of this example the DNA cassette contained the triplet TTT for codon 50 which encodes Phe. This plasmid was designated pF50. The mutant subtilisin was designated F50

B. Construction of Mutation Between Codons 122 and 127

The procedure of Example 2A was followed in substantial detail except that the mutagenesis primer of Figure 11, line 7 was used and restriction-purification for the EcoRV site in p Δ 124 was used. In addition, the DNA cassette (shaded sequence, Figure 11, line 6) contained the triplet ATT for codon 124 which encodes lie and CTT for Leu. Those plasmids which contained the substitution of lie for Met124were designeated pl124. The mutant subtilisin was designated l124.

C. Construction of Various F50/I124/Q222 Multiple Mutants

The triple mutant, F50/I124/Q222, was constructed from a three-way ligation in which each fragment contained one of the three mutations. The single mutant Q222 (pQ222) was prepared by cassette mutagenesis as described in EPO Publication No. 0130756. The F50 mutation was contained on a 2.2kb Avall to Pvull fragment from pF50; the I124 mutation was contained on a 260 bp Pvull to Avall fragment from pI124; and the Q222 mutation was contained on 2.7 kb Avall to Avall fragment from pQ222. The three fragments were ligated together and transformed into E. coli MM294 cells. Restriction analysis of plasmids from isolated transformants confirmed the construction. To analyze the final construction it was convenient that the Avall site at position 798 in the wild-type subtilisin gene was eliminated by the I124 construction.

The F50/Q222 and I124/Q222 mutants were constructed in a similar manner except that the appropriate fragment from pS4.5 was used for the final construction.

D. Oxidative Stability of Q222 Mutants

The above mutants were analyzed for stability to peracid oxidation. As shown in Fig. 12, upon incubation with diperdodecanoic acid (protein 2mg/mL, oxidant 75ppm[0]), both the I124/Q222 and the F50/I124/Q222 are completely stable whereas the F50/Q222 and the Q222 are inactivated. This indicates that conversion of Met124 to I124 in subtilisin Q222 is sufficient to confer resistance to organic peracid oxidants.

35 EXAMPLE 3

Subtilisin Mutants Having Altered Substrate Specificity-Hydrophobic Substitutions at Residues 166

Subtilisin contains an extended binding cleft which is hydrophobic in character. A conserved glycine at residue 166 was replaced with twelve non-ionic amino acids which can project their side-chains into the S-1 subsite. These mutants were constructed to determine the effect of changes in size and hydrophobicity on the binding of various substrates.

A. <u>Kinetics for Hydrolysis of Substrates Having Altered P-1 Amino Acids by Subtilisin from B.</u> Amyloliquefaciens

Wild-type subtilisin was purified from <u>B. subtilis</u> culture supernatants expressing the <u>B. amyloliquefaciens</u> subtilisin gene (Wells, J.A., et <u>al.</u> (1983) <u>Nucleic Acids Res.</u> 11, 7911-7925) as previously described (Estell, D.A., et <u>al.</u> (1985) <u>J. Biol. Chem.</u> 260, 6518-6521). Details of the synthesis of tetrapeptide substrates having the form succinyl-L-AlaL-AlaL-ProL-[X]-p-nitroanilide (where X is the P1 amino acid) are described by DelMar, E.G., et <u>al.</u> (1979) <u>Anal. Biochem.</u> 99, 316-320. Kinetic parameters, Km(M) and kcat-(s⁻¹) were measured using a modified progress curve analysis (Estell, D.A., et <u>al.</u> (1985) <u>J. Biol. Chem.</u> 260, 6518-6521). Briefly, plots of rate versus product concentration were fit to the differential form of the rate equation using a non-linear regression algorithm. Errors in kcat and Km for all values reported are less than five percent. The various substrates in Table VIII are ranged in order of decreasing hydrophobicity. Nozaki, Y. (1971), J. Biol. Chem. 246, 2211-2217; Tanford C. (1978) Science 200, 1012).

TABLE VIII

P1 substrate Amino Acid	kcat(S ⁻¹)	1/Km(M ¹)	kcat/Km (s-1M-1)
Phe	50	7,100	360,000
Tyr	28	40,000	1,100,000
Leu	24	3,100	75,000
Met	13	9,400	120,000
His	7.9	1,600	13,000
Ala	1.9	5,500	11,000
Gly	0.003	8,300	21
Gln	3.2	2,200	7,100
Ser	2.8	1,500	4,200
Glu	0.54	32	16

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The ratio of kcat/Km (also referred to as catalytic efficienty) is the apparent second order rate constant for the conversion of free enzyme plus substrate (E+S) to enzyme plus products (E+P) (Jencks, W.P., Catalysis in Chemistry and Enzymology (McGraw-Hill, 1969) pp. 321-436; Fersht, A., Enzyme Structure and Mechanism (Freeman, San Francisco, 1977) pp. 226-287). The log (kcat/Km) is proportional to transition state binding energy, ΔG_1^* . A plot of the log kcat/Km versus the hydrophobicity of the P1 side-chain (Figure 14) shows a strong correlation (r = 0.98), with the exception of the glycine substrate which shows evidence for non-productive binding. These data show that relative differences between transition-state binding energies can be accounted for by differences in P-1 side-chain hydrophobicity. When the transition-state binding energies are calculated for these substrates and plotted versus their respective side-chain hydrophobicities, the line slope is 1.2 (not shown). A slope greater than unity, as is also the case for chymotrypsin (Fersht, A., Enzyme Structure and Mechanism (Freeman, San Francisco, 1977) pp. 226-287; Harper, J.W., et al. (1984) Biochemistry, 23, 2995-3002), suggests that the P1 binding cleft is more hydrophobic than ethanol or dioxane solvents that were used to empirically determine the hydrophobicity of amino acids (Nozaki, Y., et al. J. Biol. Chem. (1971) 246, 2211-2217; Tanford, C. (1978) Science 200, 1012).

For amide hydrolysis by subtilisin, kcat can be interpreted as the acylation rate constant and Km as the dissociation constant, for the Michaelis complex (E•S), Ks. Gutfreund, H., et al (1956) Biochem. J. 63, 656. The fact that the log kcat, as well as log 1/Km, correlates with substrate hydrophobicity is consistent with proposals (Robertus, J.D., et al. (1972) Biochemistry 11, 2439-2449; Robertus, J.D., et al. (1972) Biochemistry 11, 4293-4303) that during the acylation step the P-1 side-chain moves deeper into the hydrophobic cleft as the substrate advances from the Michaelis complex (E•S) to the tetrahedral transition-state complex (E•S*). However, these data can also be interpreted as the hydrophobicity of the P1 side-chain effecting the orientation, and thus the susceptibility of the scissile peptide bond to nucleophilic attack by the hydroxyl group of the catalytic Ser221.

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The dependence of kcat/Km on P-1 side chain hydrophobicity suggested that the kcat/Km for hydrophobic substrates may be increased by increasing the hydrophobicity of the S-1 binding subsite. To test this hypothesis, hydrophobic amino acid substitutions of Gly166 were produced.

Since hydrophobicity of aliphatic side-chains is directly proportional to side-chain surface area (Rose, G.D., et al. (1985) Science 229, 834-838; Reynolds, J.A., et al. (1974) Proc. Natl. Acad. Sci. USA 71, 2825-2927), increasing the hydrophobicity in the S-1 subsite may also sterically hinder binding of larger substrates. Because of difficulties in predicting the relative importance of these two opposing effects, we elected to generate twelve non-charged mutations at position 166 to determine the resulting specificities against non-charged substrates of varied size and hydrophobicity.

B. Cassette Mutagenesis of the P1 Binding Cleft

The preparation of mutant subtilisims containing the substitution of the hydrophobic amino acids Ala, Val and Phe into residue 166 has been described in EPO Publication No. 0130756. The same method was used to produce the remaining hydrophobic mutants at residue 166. In applying this method, two unique and silent restriction sites were introduced in the subtilisin genes to closely flank the target codon 166. As can be seen in Figure 13, the wild type sequence (line 1) was altered by site-directed mutagenesis in M13 using the indicated 37mer mutagenesis primer, to introduce a 13 bp delection (dashedline) and unique Sacl and Xmal sites (underlined sequences) that closely flank codon 166. The subtilisin gene fragment was subcloned back into the E. coli - B. subtilis shuttle plasmid, pBS42, giving the plasmid pΔ166 (Figure 13,

line 2). p∆166 was cut open with SacI and XmaI, and gapped linear molecules were purified (Figure 13, line 3). Pools of synthetic oligonucleotides containing the mutation of interest were annealed to give duplex DNA cassettes that were ligated into gapped p∆166 (underlined and overlined sequences in Figure 13, line 4). This construction restored the coding sequence except over position 166(NNN; line 4). Mutant sequences were confirmed by dideoxy sequencing. Asterisks denote sequence changes from the wild type sequence. Plasmids containing each mutant B. amyloliquefaciens subtilisin gene were expressed at roughly equivalent levels in a protease deficient strain of B. subtilis, BG2036 as previously described. EPO Publication No. 0130756; Yang, M., et al. (1984) J. Bacteriol. 160, 15-21; Estell, D.A., et al (1985) J. Biol. Chem. 260, 6518-6521.

C. Narrowing Substrate Specificity by Steric Hindrance

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To probe the change in substrate specificity caused by steric alterations in the S-1 subsite, position 166 mutants were kinetically analyzed versus P1 substrates of increasing size (i.e., Ala, Met, Phe and Tyr). Ratios of kcat/Km are presented in log form in Figure 15 to allow direct comparisons of transition-state binding energies between various enzyme-substrate pairs.

According to transition state theory, the free enery difference between the free enzyme plus substrate (E + S) and the transition state complex $(E \cdot S^*)$ can be calculated from equation (1),

(1)
$$^{\Delta}G_{T}^{\neq}$$
 = -RT ln kcat/Km + RT ln kT/h

in which kcat is the turnover number, Km is the Michaelis constant, R is the gas constant, T is the temperature, k is Boltzmann's constant, and h is Planck's constant. Specificity differences are ezpressed quantitatively as differences between transition state binding energies (i.e., ΔΔG^{*}_t), and can be calculated from equation (2).

(2)
$$^{\Delta\Delta}G_{T}^{\neq} = -RT \ln (kcat/Km)_{A}/(kcat/Km)_{B}$$

35 A and B represent either two different substrates assayed againt the same enzyme, or two mutant enzymes assayed against the same substrate.

As can be seen from Figure 15A, as the size of the side-chain at position 166 increases the substrate preference shifts from large to small P-1 side-chains. Enlarging the side-chain at position 166 causes kcat/Km to decrease in proportion to the size of the P-1 substrate side-chain (e.g., from Gly166 (wild-type) through W166, the kcat/Km for the Tyr substrate is decreased most followed in order by the Phe, Met and Ala P-1 substrates).

Specific steric changes in the position 166 side-chain, such as he presence of a β -hydroxyl group, β - or γ -aliphatic branching, cause large decreases in kcat/Km for larger P1 substrates. Introducing a β -hydroxyl group in going from A166 (Figure 15A) to S166 (Figure 15B), causes an 8 fold and 4 fold reduction in kcat/Km for Phe and Tyr substrates, respectively, while the values for Ala and Met substrates are unchanged. Producing a β -branched structure, in going from S166 to T166, results in a drop of 14 and 4 fold in kcat/Km for Phe and Tyr, respectively. These differences are slightly magnified for V166 which is slightly larger and isosteric with T166. Enlarging the β -branched substituents from V166 to I166 causes a lowering of kcat/Km between two and six fold toward Met, Phe and Tyr substrates. Inserting a γ -branched structure, by replacing M166 (Figure 15A) with L166 (Figure 15B), produces a 5 fold and 18 fold decrease in kcat/Km for Phe and Tyr substrates, respectively. Aliphatic γ -branched appears to induce less steric hindrance toward the Phe P-1 substrate than β -branching, as evidenced by the 100 fold decrease in kcat/Km for the Phe substrate in going from L166 to I166.

Reductions in kcat/Km resulting from increases in side chain size in the S-1 subsite, or specific structural features such as β - and γ -branching, are quantitatively illustrated in Figure 16. The kcat/Km values for the position 166 mutants determined for the Ala, Met, Phe, and Tyr P-1 substrates (top panel through bottom panel, respectively), are plotted versus the position 166 side-chain volumes (Chothia, C. (1984) Ann. Rev. Biochem. 53, 537-572). Catalytic efficiency for the Ala substrate reaches a maximum for

I166, and for the Met substrate it reaches a maximum between V166 and L166. The Phe substrate shows a broad kcat/Km peak but is optimal with A166. Here, the β -branched position 166 substitutions form a line that is parallel to, but roughly 50 fold lower in kcat/Km than side-chains of similar size (i.e., C166 versus T166, L166 versus I166). The Tyr substrate is most efficiently utilized by wild type enzyme (Gly166), and there is a steady decrease as one proceeds to large position 166 side-chains. The β -branched and γ -branched substitutions form a parallel line below the other non-charged substitutions of similar molecular volume.

The optimal substitution at position 166 decreases in volume with increasing volume of the P1 substrate [i.e., I166/Ala substrate, L166/Met substrate, A166/Phe substrate, Gly166/Tyr substrate]. The combined volumes for these optimal pairs may approximate the volume for productive binding in the S-1 subsite. For the optimal pairs, Gly166/Tyr substrate, A166/Phe substrate, L166/Met substrate, V166/Met substrate, and I166/Ala substrate, the combined volumes are 266,295,313,339 and 261 A³, respectively. Subtracting the volume of the peptide backbone from each pair (i.e., two times the volume of glycine), an average sidechain volume of 160±32A³ for productive binding can be calculated.

The effect of volume, in excess to the productive binding volume, on the drop in transition-state binding energy can be estimated from the Tyr substrate curve (bottom panel, Figure 16), because these data, and modeling studies (Figure 2), suggest that any substitution beyond glycine causes steric repulsion. A best-fit line drawn to all the data (r = 0.87) gives a slope indicating a loss of roughly 3 kcal/mol in transition state binding energy per $100A^3$ of excess volume. ($100A^3$ is approximately the size of a leucyl side-chain.)

D. Enhanced Catalytic Efficiency Correlates with Increasing Hydrophobicity of the Position 166 Substitution

Substantial increases in kcat/Km occur with enlargement of the position 166 side-chain, except for the Tyr P-1 substrate (Figure 16). For example, kcat/Km increases in progressing from Gly166 to I166 for the Ala substrate (net of ten-fold), from Gly166 to L166 for the Met substrate (net of ten-fold) and from Gly166 to A166 for the Phe substrate (net of two-fold). The increases in kcat/Km cannot be entirely explained by the attractive terms in the van der Waals potential energy function because of their strong distance dependence (1/r⁶) and because of the weak nature of these attractive forces (Jencks, W.P., Catalysis in Chemistry and Enzymology (McGraw-Hill, 1969) pp. 321-436; Fersht, A., Enzyme Structure and Mechanism (Freeman, San Francisco, 1977) pp. 226-287; Levitt, M. (1976) J. Mol. Biol. 104, 59-107). For example, Levitt (Levitt, M. (1976) J. Mol. Biol. 104, 59-107) has calculated that the van der Waals attraction between two methionyl residues would produce a maximal interaction energy of roughly -0.2 kcal/mol. This energy would translate to only 1.4 fold increase in kcat/Km.

The increases of catalytic efficiency caused by side-chain substitutions at position 166 are better accounted for by increases in the hydrophobicity of the S-1 subsite. The increase kcat/Km observed for the Ala and Met substrates with increasing position 166 side-chain size would be expected, because hydrophobicity is roughly proportional to side-chain surface area (Rose, G.D., et al. (1985) Science 229, 834-838; Reynolds, J.A., et al. (1974) Proc. Natl. Acad. Sci. USA 71, 2825-2927).

Another example that can be interpreted as a hydrophobic effect is seen when comparing kcat/Km for isosteric substitutions that differ in hydrophobicity such as S166 and C166 (Figure 16). Cysteine is considerably more hydrophobic than serine (-1.0 versus +0.3 kcal/mol) (Nozaki, Y., et al. (1971) J. Biol. Chem. 246, 2211-2217; Tanford, C. (1978) Science 200, 1012). The difference in hydrophobicity correlates with the observation that C166 becomes more efficient relative to Ser166 as the hydrophobicity of the substrates increases (i.e., Ala < Met < Tye < Phe). Steric hindrance cannot explain these differences because serine is considerably smaller than cysteine (99 versus 118A³). Paul, I.C., Chemistry of the -SH Group (ed. S. Patai, Wiley Interscience, New York, 1974) pp. 111-149.

E. Production of an Elastase-Like Specificity in Subtilisin

The I166 mutation illustrates particularly well that large changes in specificity can be produced by altering the structure and hydrophobicity of the S-1 subsite by a single mutation (Figure 17). Progressing through the small hydrophobic substrates, a maximal specificity improvement over wild type occurs for the Val substrate (16 fold in kcat/Km). As the substrate side chain size increases, these enhancements shrink to near unity (i.e., Leu and His substrates). The I166 enzyme becomes poorer against larger aromatic substrates of increasing size (e.g., I166 is over 1,000 fold worse against the Tyr substrate than is Gly166). We interpret the increase in catalytic efficiency toward the small hydrophobic substrates for I166 compared to Gly166 to the greater hydrophobicity of isoluecine (i.e., -1.8 kcal/mol versus 0). Nozaki, Y., et al. (1971) J. Biol. Chem. 246, 2211-2217; Tanford, C. (1978) Science 200, 1012. The decrease in catalytic efficiency

toward the very large substrates for I166 versus Gly166 is attributed to steric repulsion.

The specificity differences between Gly166 and I166 are similar to the specificity differences between chymotrypsin and the evolutionary relative, elastase (Harper, J.W., et al (1984) Biochemistry 23, 2995-3002). In elastase, the bulky amino acids, Thr and Val, block access to the P-1 binding site for large hydrophobic substrates that are preferred by chymotrypsin. In addition, the catalytic efficiencies toward small hydrophobic substrates are greater for elastase than for chymotrypsin as we obeseve for I166 versus Gly166 in subtilisin.

EXAMPLE 4

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Substitution of Ionic Amino Acids for Gly166

The construction of subtilisin mutants containing the substitution of the ionic amino acids Asp, Asn, Gln, Lys and Ang are disclosed in EPO Publication No. 0130756. The present example describes the construction of the mutant subtilisin containing Glu at position 166 (E166) and presents substrate specificity data on these mutants. Further data on position 166 and 156 single and double mutants is presented infra.

pΔ166, described in Example 3, was digested with SacI and XmaI. The double strand DNA cassette (underlined and overlined) of line 4 in Figure 13 contained the triplet GAA for the codon 166 to encode the replacement of Glu for Gly166. This mutant plasmid designated pQ166 was propagated in BG2036 as described. This mutant subtilisin, together with the other mutants containing ionic substituent amino acids at residue 166, were isolated as described and further analyzed for variations in substrate specificity.

Each of these mutants was analyzed with the tetrapeptide substrates, succinyl-L-AlaL-AlaProL-X-p-nitroanilide, where X was Phe, Ala and Glu.

The results of this analysis are shown in Table IX.

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TABLE IX

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Position 166	P-1 Su	bstrate (kcat/	Km x 10 ⁻⁴)
	Phe	Ala	Glu
Gly (wild type)	36.0	1.4	0.002
Asp (D)	0.5	0.4	<0.001
Glu (E)	3.5	0.4	<0.001
Asn (N)	18.0	1.2	0.004
Gin (Q)	57.0	2.6	0.002
Lys (K)	52.0	2.8	1.2
Arg (R)	42.0	5.0	80.0

These results indicate that charged amino acid substitutions at Gly166 have improved catalytic efficiencies (kcat/Km) for oppositely charged P-1 substrates (as much as 500 fold) and poorer catalytic efficiency for like charged P-1 substrates.

EXAMPLE 5

Substitution of Glycine at Position 169

The substitution of Gly169 in B. <u>amyloliquefacions</u> subtilisin with Ala and Ser is described in EPO Publication No. 0130756. The same method was used to make the remaining 17 mutants containing all other substituent amino acids for position 169.

The construction protocol is summarized in Figure 18. The overscored and underscored double stranded DNA cassettes used contained the following triplet encoding the substitution of the indicated amino acid at residue 169.

GCT	Α	ATG	М
TGT	С	AAC	N
GAT	D	CCT	Р
GAA	Е	CAA	Q
тс	F	AGA	R
GGC	G	AGC	S
CAC	н	ACA	Т
ATC	1	GTT	٧
AAA	Κ	TGG	W
CTT	L	TAC	Υ

Each of the plasmids containing a substituted Gly169 was designated pX169, where X represents the substituent amino acid. The mutant subtilisins were similarly designated.

Two of the above mutant subtilisins, A169 and S169, were analyzed for substrate specificity against synthetic substrates containing Phe, Leu, Ala and Arg in the P-1 position. The following results are shown in Table X.

TABLE X

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Effect of Serine and Al	anine Mutations	at Position 169	on P-1 Substra	ate Specificity
Position 169		P-1 Substrate [l	cat/Km x 10	⁴)
	Phe	Leu	Ala	Arg
Gly (wild type)	40	10	1	0.4
A169	120	20	1	0.9
S169	50	10	1	0.6

These results indicate that substitutions of Ala and Ser at Gly169 have remarkably similar catalytic efficiencies against a range of P-1 substrates compared to their position 166 counterparts. This is probably because position 169 is at the bottom of the P-1 specificity subsite.

EXAMPLE 6

Substitution at Position 104

Tyr104 has been substituted with Ala, His, Leu, Met and Ser. The method used was a modification of the site directed mutagenesis method. According to the protocol of Figure 19, a primer (shaded in line 4) introduced a unique HindIII site and a frame shift mutation at codon 104. Restriction-purification for the unique HindIII site facilitated the isolation of the mutant sequence (line 4). Restriction-selection against this HindIII site using pimers in line 5 was used to obtain position 104 mutants.

The following triplets were used in the primers of Figure 19, line 5 for the 104 codon which substituted the following amino acids.

GCT TTC Р **ATG** М CCT **ACA** Т CTT L AGC S TGG W CAC Н TAC Q **GTT** ٧ CAA Ε R GAA AGA GGC G Ν AAC D **ATC** ı GAT κ Ç AAA **TGT**

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The substrates in Table XI were used to analyze the substrate specificity of these mutants. The results obtained to H104 subtilisin are shown in Table XI.

TABLE XI

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Substrate	ko	at	к	(m	Kca	ıt/Km
	WT	H104	WT	H104	WT	H104
sAAPFpNA	50.0	22.0	1.4x10 ⁻⁴	7.1x10 ⁻⁴	3.6x10 ⁵	3.1x10 ⁴
sAAPApNA	3.2	2.0	2.3x10 ⁻⁴	1.9x10 ⁻³	1.4x10⁴	1×10³
sFAPFpNA	26.0	38.0	1.8x10 ⁻⁴	4.1x10 ⁻⁴	1.5x10 ⁵	9.1x10⁴
sFAPApNA	0.32	2.4	7.3x10 ⁻⁵	1.5x10 ⁻⁴	4.4x10 ³	1.6x10 ⁴

From these data it is clear that the substitution of His for Tyr at position 104 produces an enzyme which is more efficient (higher kcat/Km) when Phe is at the P-4 substrate position than when Ala is at the P-4 substrate position.

EXAMPLE 7

Substitution of Ala152

Ala152 has been substituted by Gly and Ser to determine the effect of such substitutions on substrate specificity.

The wild type DNA sequence was mutated by the V152/P153 primer (Figure 20, line 4) using the above restriction-purification approach for the new <u>KpnI</u> site. Other mutant primers (shaded sequences Figure 20; S152, line 5 and G152, line 6) mutated the new <u>KpnI</u> site away and such mutants were isolated using the restriction-selection procedure as described above for loss of the KpnI site.

The results of these substitutions for the above synthetic substrates containing the P-1 amino acids Phe, Leu and Ala are shown in Table XII.

TABLE XII

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P-1 Su	bstrate (kcat/l	(mx10 ⁻⁴)
Phe	Leu	Ala
0.2 40.0	0.4 10.0	<0.04 1.0 0.2
	Phe 0.2	0.2 0.4 40.0 10.0

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These results indicate that, in contrast to positions 166 and 169, replacement of Ala152 with Ser or Gly causes a dramatic reduction in catalytic efficiencies across all substrates tested. This suggests Ala152, at the top of the S-1 subsite, may be the optimal amino acid because Ser end Gly ore homologous Ala substitutes.

EXAMPLE 8

Substitution at Position 156

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Mutants containing the substitution of Ser and Gln for Glu156 have been constructed according to the overall method depicted in Figure 21. This method was designed to facilitate the construction of multiple mutants at position 156 and 166 as will be described hereinafter. However, by regenerating the wild type Gly166, single mutations at Glu156 were obtained.

The plasmid $p\Delta 166$ is already depicted in line 2 of Figure 13. The synthetic oligonucleotides at the top right of Figure 21 represent the same DNA cassettes depicted in line 4 of Figure 13. The plasmid p166 in Figure 21 thus represents the mutant plasmids of Examples 3 and 4. In this particular example, p166 contains the wild type Gly166.

Construction of position 156 single mutants were prepared by ligation of the three fragments (1-3) indicated at the bottom of Figure 21. Fragment 3, containing the carboxy-terminal portion of the subtilisin gene including the wild type position 166 codon, was isolated as a 610 bp Sacl-BamHI fragment. Fragment 1 contained the vector sequences, as well as the amino-terminal sequences of the subtilisin gene through codon 151. To produce fragment 1, a unique Kpnl site at codon 152 was introduced into the wild type subtilisin sequence from pS4.5. Site-directed mutagenesis in M13 employed a primer having the sequence 5'-TA-GTC-GTT-GCG-GTA-CCC-GGT-AAC-GAA-3' to produce the mutation. Enrichment for the mutant sequence was accomplished by restriction with KpnI, purification and self ligation. The mutant sequence containing the KpnI site was confirmed by direct plasmid sequencing to give pV152. pV152 (~1 µg) was digested with Kpnl and treated with 2 units of DNA polymerase I large fragment (Klenow fragment from Boeringer-Mannheim) plus 50 µM deoxynucleotide triphosphates at 37 °C for 30 min. This created a blunt end that terminated with codon 151. The DNA was extracted with 1:1 volumes phenol and CHCl3 and DNA in the aqueous phase was precipitated by addition of 0.1 volumes 5M ammonium acetate and two volumes ethanol. After centrifugation and washing the DNA pellet with 70% ethanol, the DNA was lyophilized. DNA was digested with BamHI and the 4.6kb piece (fragment 1) was purified by acrylamide gel electrophoresis followed by electroelution. Fragment 2 was a duplex synthetic DNA cassette which when ligated with fragments 1 and 3 properly restored the coding sequence except at codon 156. The top strand was synthesized to contain a glutamine codon, and the complementary bottom strand coded for serine at 156. Ligation of heterophosphorylated cassettes leads to a large and favorable bias for the phosphorylated over the non-phosphorylated oligonucleotide sequence in the final segrated plasmid product. Therefore, to obtain Q156 the top strand was phosphorylated, and annealed to the non-phosphorylated bottom strand prior to ligation. Similarly, to obtain \$156 the bottom strand was phosphorylated and annealed to the nonphosphorylated top strand. Mutant sequences were isolated after ligation and transformation, and were confirmed by restriction analysis and DNA sequencing as before. To express variant subtilisins, plasmids were transformed into a subtilisin-neutral protease deletion mutant of B. subtilis, BG2036, as previously described. Cultures were fermented in shake flasks for 24 h at 37°C in LB media containing 12.5 mg/mL chloramphenicol and subtilisin was purified from culture supernatants as described. Purity of subtilisin was greater than 95% as judged by SDS PAGE.

These mutant plasmids designated pS156 and pQ156 and mutant subtilisins designated S158 and Q156 were analyzed with the above synthetic substrates where P-1 comprised the amino acids Glu, Gln, Met and Lys. The results of this analyses are presented in Example 9.

EXAMPLE 9

Multiple Mutants With Attered Substrate Specificity - Substitution at Positions 156 and 166

Single substitutions of position 166 are described in Examples 3 and 4. Example 8 describes single substitutions at position 156 as well as the protocol of Figure 21 whereby various double mutants comprising the substitution of various amino acids at positions 156 and 166 can be made. This example describes the construction and substrate specificity of subtilisin containing substitutions at position 156 and 166 and summarizes some of the data for single and double mutants at positions 156 and 166 with various substrates.

K166 is a common replacement amino acid in the 156/166 mutants described herein. The replacement of Lys for Gly166 was achieved by using the synthetic DNA cassette at the top right of Figure 21 which contained the triplet AAA for NNN. This produced fragment 2 with Lys substituting for Gly166.

The 156 substituents were Gln and Ser. The Gln and Ser substitutions at Gly156 are contained within fragment 3 (bottom right Figure 21).

The multiple mutants were produced by combining fragments 1, 2 and 3 as described in Example 8. The mutants Q156/K168 and S156/K166 were selectively generated by differential phosphorylation as described. Alternatively, the double 156/166 mutants, c.f. Q156/K168 and S156/K166, were prepared by ligation of the 4.6kb Sacl-BamHI fragment from the relevant p156 plasmid containing the 0.6kb Sacl-BamHI fragment from the relevant p166 plasmid.

These mutants, the single mutant K166, and the S156 and Q156 mutants of Example 8 were analyzed for substitute specificity against synthetic polypeptides containing Phe or Glu as the P-1 substrate residue. The results are presented in Table XIII.

5		kcat/Km (mutant) kcat/Km(wt)	(1)	(1)	1.4	750	4.4	3100	4.4	1000	2.0	6.9	3.1	17
15		kcat/Km	3.6×10 ⁵	1.6×10^{1}	5.2×10 ⁵	1.2×10 ⁴	1.6×10 ⁶	5.0×10 ⁴	1.6×10 ⁶	1.6×10 ⁴	7.3×10 ⁵	1.1×10 ²	1.1×10 ⁶	2.7×10 ²
20		1	4	~ :	ıcı ı	വ	ru	ري د	ហ	S	Ŋ	m	S.	6 0
25	TABLE XIII	Km	1.4×10-4	3.4×10	4.0×10 ⁻	5.6×10	1.9×10 ⁻	3.1×10 ⁻	1.8×10	3.9×10 ⁻	4.7×10	1.8×10	4.5×10	3.3×10 ⁻³
30	TAI	kcat	50.00	0.54	20.00	0.70	30.00	1.60	30.00	09.0	34.00	0.40	48.00	06.0
35		Substrate P-1 Residue	Phe	Glu	Phe	Glu	Phe	Glu	Phe	Glu	Phe	Glu	Phe	Glu
40		(q) Pa	WT.)											
45		Enzymes Compare	Glu156/Gly166 (WT)		95		Q156/R166		S156/K166		99		99	
50		គ្ន	G1 ^u		K166		01!		\$15		S156		E156	

As can be seen in Table XIV, either of these single mutations improve enzyme performance upon substrates with glutamate at the P-1 enzyme binding site. When these single mutations were combined, the resulting multiple enzyme mutants are better than either parent. These single or multiple mutations also alter the relative pH activity profiles of the enzymes as shown in Figure 23.

To isolate the contribution of electrostatics to substrate specificity from other chemical binding forces, these various single and double mutants were analyzed for their ability to bind and cleave synthetic substrates containing Glu, Gln, Met and Lys as the P-1 substrate amino acid. This permitted comparisons between side-chains that were more sterically similar but differed in charge (e.g., Glu versus Gln, Lys versus Met). Similarly, mutant enzymes were assayed against homologous P-1 substrates that were most sterically similar but differed in charge (Table XIV).

10				Lys		(3.00)	(3.69)	(2.88)	(3.15)	(3.22)	(3.07)	(3.89)	(3.24)	(3.13)	(2.82)	(2.74)	(2.74)	(2.80)	(2.80)	(2.93)	(2.75)	(2.84)		(-1.0)
15			/Km) (c)	1		4.23	4.48	4.15	4.10	4.41	4.24	4.70	4.90	4.60	3.76	3.46	3.75	3.68	3.19	4.23	3.23	3.73		-1.3
20			kcat/Km (log 1/Km) (c)	Met		(2.74)	(3.28)	(3.82)	(4.36)	(3.87)	(3.68)	(4.83)	(4.46)	(3.97)	(4.61)	(4.55)	(4.66)	(4.64)	(4.22)	(4.45)	(4.68)	(4.90)		(2.2)
		Subtilisins Substrates	kcat/Km	Σ		3.93	3.86	4.99	5.43	4.94	4.67	5.64	5 ; 65	5,07	5:77	5.61	5.79	5.72	5.32	6.15	5.97	6.16		2.3
25			log			(2.56)	(2.91)	(3.14)	(3.64)	(3.08)	(3.09)	(3.19)	(3.55)	(3:35)	(3.81)	(3.68)	(3.76)	(3.82)	(3.50)	(3.88)	(3.68)	(3.94)		(1.4)
30	XIV	tion 156/166 Different Pl	Substrate	S		3.02	3.06	3.85	4.36	3.40	3.41	3.89	4.34	3.85	4.53	4.09	4.51	4.57	4.26	4.70	4.64	4.84		1.8
35	TABLE	Position for Diffe	P-1	Glu				(2.22)	(2.12)	(1.79)	(2.13)	(2.30)		(1.47)	(2.48)	(2.73)	(2.72)	(2.78)	(3.30)	(4.25)	(4.50)	(4.40)		(3.0)
40		o f ned		S 		n.d.	n.d.	. 1.62	1.20	1.30	1.23	1.20	n.d.	1.20	2.42	2.31	2.04	1.91	2.91	4.09	4.70	4.21		3.5
		Kinetics Determi		<u> </u>																٠			-	ē
45			Net	Charge		-2	-2	7	7	-1	-1	-1	-1	7	0	0	0	0	0	0	+1	+1	ence:	J 1/Km) (d)
50				a)										wt)									Maximum differen	log kcat/Km (log
			уте,	Position (a)	166	Asp	Glu	Asn	Gln	Asp	Asp	Met	Ala	Gly(wt)	G1y	Gly	Asn	Asn	Arg	Lys	Lys	Lys	mum d	kcat/
55			Enz	Posi	156	Glu	Glu	Glu	Glu	Gln	Ser	Glu	Glu	Glu	Gln	Ser	Gln	Ser	Gla	Glu	Gln	Ser	Maxi	109

Footnotes to Table XIV:

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- (a) B. <u>subtilis</u>, BG 2036, expressing indicated variant subtilisin were fermented and enzymes purified as previously described (Estell, <u>et al</u>. (1985) <u>J. Biol. Chem. 260</u>, 6518-6521). Wild type subtilisin is indicated (wt) containing Glul56 and Glyl66.
- (b) Net charge in the P-1 binding site is defined as the sum of charges from positions 156 and 166 at pH 8.6.
- (C) Values for kcat(s⁻¹) and Km(M) were measured in 0.1M Tris pH 8.6 at 25°C as previously described · P-1 substrates having the form succinyl-L-AlaL-AlaL-ProL-[X]-p-nitroanilide, where X is the indicated P-1 amino acid. Values for log 1/Km parentheses. shown inside All errors determination of kcat/Km and 1/Km are below 5%.
- (d) Because values for Glul56/Aspl66(Dl66) are too small to determine accurately, the maximum difference taken for GluP-1 substrate is limited to a charge range of +1 to -1 charge change.

n.d. = not determined

The kcat/Km ratios shown are the second order rate constants for the conversion of substrate to product, and represent the catalytic efficiency of the enzyme. These ratios are presented in logarithmic form to scale the data, and because log kcat/Km is proportional to the lowering of transition-state activation energy (ΔG_T). Mutations at position 156 and 166 produce changes in catalytic efficiency toward Glu, Gln, Met and Lys P-1 substrates of 3100, 60, 200 and 20 fold, respectively. Making the P-1 binding-site more positively charged [e.g., compare Gln156/Lys166 (Q156/K166) versus Glu156/Met166 (Glu156/M166)] dramatically increased kcat/Km toward the Glu P-1 substrate (up to 3100 fold), and decreased the catalytic efficiency toward the Lys P-1 substrate (up to 10 fold). In addition, the results show that the catalytic efficiency of wild type enzyme can be greatly improved toward any of the four P-1 substrates by mutagenesis of the P-1 binding site.

The changes in kcat/Km ore caused predominantly by changes in 1/Km. Because 1/Km is approximately equal to 1/Ks, the enzyme-substrate association constant, the mutations primarily cause a change in substrate binding. These mutations produce smaller effects on kcat that run parallel to the effects on 1/Km. The changes in kcat suggest either an alteration in binding in the P-1 binding site in going from the Michaelis-complex E+S) to the transition-state complex (E-S#) as previously proposed (Robertus, J.D., et al. (1972) Biochemistry 11, 2439-2449; Robertus, J.D., et al. (1972) Biochemistry 11, 4293-4303), or change in the position of the scissile peptide bond over the catalytic serine in the E+S complex.

Changes in substrate preference that arise from changes in the net charge in the P-1 binding site show trends that are best accounted for by electrostatic effects (Figure 28). As the P-1 binding cleft becomes more positively charged, the average catalytic efficiency increases much more for the Glu P-1 substrate than for its neutral and isosteric P-1 homolog, Gln (Figure 28A). Furthermore, at the positive extreme both substrates have nearly identical catalytic efficiencies.

In contrast, as the P-1 site becomes more positively charged the catalytic efficiency toward the Lys P-1 substrate decreases, and diverges sharply from its neutral and isosteric homolog, Met (Figure 28B). The similar and parallel upward trend seen with increasing positive charge for the Met and Glu P-1 substrates probably results from the fact that all the substrates are succinylated on their amino-terminal end, and thus carry a formal negative charge.

The trends observed in log kcat/Km are dominated by changes in the Km term (Figures 28C and 28D). As the pocket becomes more positively charged, the log 1/Km values converge for Glu and Gln P-1 substrates (Figure 28C), and diverge for Lys and Met P-1 substrates (Figure 28D). Although less

pronounced effects are seen in log kcat, the effects of P-1 charge on log kcat parallel those seen in log 1/Km and become larger as the P-1 pocket becomes more positively charged. This may result from the fact that the transition-state is a tetrahedral anion, and a net positive charge in the enzyme may serve to provide some added stabilization to the transition-state.

The effect of the change in P-1 binding-site charge on substrate preference can be estimated from the differences in slopes between the charged and neutral isosteric P-1 substrates (Figure 28B). The average change in substrate preference (Δlog kcat/Km) between charged and neutral isosteric substrates increases roughly 10-fold as the complementary charge or the enzyme increases (Table XV). When comparing Glu versus Lys, this difference is 100-fold and the change in substrate preference appears predominantly in the Km term.

TABLE XV

15	Differential Effect on Binding Site Charge on log kcat/Km or (log Charge ^(a)	1/Km) for P-1	Substrates	that Differ in
	Change in P-1 Binding Site Charge ^(b)	∆log ko	at/Km (∆lo	g 1/Km)
		GluGln	MetLys	GluLys
20	-2 to -1 -1 to 0 0 to +1 Avg. change in log kcat/K _m or (log 1/Km) per unit charge change	n.d. 0.7 (0.6) 1.5 (1.3) 1.1 (1.0)	1.2 (1.2) 1.3 (0.8) 0.5 (0.3) 1.0 (0.8)	n.d. 2.1 (1.4) 2.0 (1.5) 2.1 (1.5)

(a) The difference in the slopes of curves were taken between the P-1 substrates over the charge interval given for log (kcat/Km) (Figure 28A, B) and (log 1/Km) (Figure 28C, D). Values represent the differential effect a charge change has in distinguishing the substrates that are compared.

(b) Charge in P-1 binding site is defined as the sum of charges from positions 156 and 166.

The free energy of electrostatic interactions in the structure and energetics of salt-bridge formation depends on the distance between the charges and the microscopic dielectric of the media. To dissect these structural and microenvironmental effects, the energies involved in specific salt-bridges were evaluated. In addition to the possible salt-bridges shown (Figures 29A and 29B), reasonable salt-bridges can be built between a Lys P-1 substrate and Asp at position 166, and between a Glu P-1 substrate and a Lys at position 166 (not shown). Although only one of these structures is confirmed by X-ray crystalography (Poulos, T.L., et al. (1976) J. Mol. Biol. 257 1097-1103), all models have favorable torsion angles (Sielecki, A.R., et al. (1979) J. Mol. Biol. 134, 781-804), and do not introduce unfavorable van der Waals contacts.

The change in charged P-1 substrate preference brought about by formation of the model salt-bridges above are shown in Table XVI.

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XVI
TABLE

Effect of Salt Bridge Formation Between Enzyme and Substrate on Pl Substrate Preference (a)

Enzymes Compared (b)	ompared (b)	Enzyme Position Changed	P-1 Substrates Compared	Substrate (d) Preference	Substrate (d) Preference og (kcat/Km)	in Substrate Preference Aalog (kcat/Km) (1-2)
G1u156/Asp166	Gln156/Asp166	156	LysMet	+0.30	-0.53	0.83
Glu156/Asn166	Gln156/Asn166	156	LysMet	-0.84	-2.04	1.20
Glu156/Gly166	Gln156/Gly166	156	LysMet	-0.47	-2.10	1.63
Glu156/Lsy-166	Gln156/Lys166	156	LysMet	-1.92	-2.74	0.82
G1u156/Asp166	G1u156/Asn166	166	LysMet	Ave AA1+0.30	og (kcat/ -0.84	Ave &&log (kcat/Km) 1.10 ± 0.3 +0.30 -0.84 1.14
G1u156/G1u166	Glu156/Glu166	166	LysMet	+0.62	-1.33	1.95
Gln156/Asp166	Gln156/Asn166	166	LysMet	-0.53	-2.04	1.51
Ser156/Asp166	Ser156/Asn166	166	LysMet	-0.43	-2.04	1.61
Glu156/Lys166	Glu156/Met166	166	GluGln	-0.63	-2.69	2/06

Footnotes to Table XVI:

- (a) Molecular modeling shows it is possibl to form a salt bridge between the indicated charged P-l substrate and a complementary charge in the P-l binding site of the enzyme at the indicated position changed.
- (b) Enzymes compared have sterically similar amino acid substitutions that differ in charge at the indicated position.
 - (c) The P-1 substrates compared are structurally similar but differ in charge. The charged P-1 substrate is complementary to the charge change at the position indicated between enzymes 1 and 2.
 - (d) Date from Table XIV was used to compute the difference in log (kcat/Km) between the charged and the non-charged P-1 substrate (i.e., the substrate preference). The substrate preference is shown separately for enzyme 1 and 2.
- (e) The difference in substrate preference between enzyme 1 (more highly charged) and enzyme 2 (more neutral) represents the rate change accompanying the electrostatic interaction.

The difference between catalytic efficiencies (i.e., Δlog kcat/Km) for the charged and neutral P-1 substrates (e.g., Lys minus Met or Glu minus Gln) give the substrate preference for each enzyme. The change in substrate preference (ΔΔlog kcat/Km) between the charged and more neutral enzyme homologs (e.g., Glu156/Gly166 minus Gln156(Q156)/Gly166) reflects the change in catalytic efficiency that may be attributed solely to electrostatic effects.

These results show that the average change in substrate preference is considerably greater when electrostatic substitutions are produced at position 166 (50-fold in kcat/Km) versus position 156 (12-fold in kcat/Km). From these $\Delta\Delta$ log kcat/Km values, an average change in transition-state stabilization energy can be calculated of -1.5 and -2.4 kcal/mol for substitutions at positions 156 and 166, respectively. This should represent the stabilization energy contributed from a favorable electrostatic interaction for the binding of free enzyme and substrate to form the transition-state complex.

EXAMPLE 10

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45 Substitutions at Position 217

Tyr217 has been substituted by all other 19 amino acids. Cassette mutagenesis as described in EPO publication No. 0130756 was used according to the protocol of Figure 22. The EcoRV restriction site was used for restriction-purification of p Δ 217.

Since this position is involved in substrate binding, mutations here effect kinetic parameters of the enzyme. An example is the substitution of Leu for Tyr at position 217. For the substrate sAAPFpNa, this mutant has a kcat of 277 5' and a Km of 4.7x10⁻⁴ with a kcat/Km ratio of 6x10⁵. This represents a 5.5-fold increase in kcat with a 3-fold increase in Km over the wild type enzyme.

In addition, replacement of Tyr217 by Lys, Arg, Phe or Leu results in mutant enzymes which are more stable at pHs of about 9-11 than the WT enzyme. Conversely, replacement of Tyr217 by Asp, Glu, Gly or Pro results in enzymes which are less stable at pHs of about 9-11 than the WT enzyme.

EXAMPLE 11

Multiple Mutants Having Altered Thermal Stability

B. amyloliquefacien subtilisin does not contain any cysteine residues. Thus, any attempt to produce thermal stability by Cys cross-linkage required the substitution of more than one amino acid in subtilisin with Cys. The following subtilisin residues were multiply substituted with cysteine:

Thr22/Ser87

Ser24/Ser87

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Mutagenesis of Ser24 to Cys was carried out with a 5' phosphorylated oligonucleotide primer having the sequence

$$5'-pC-TAC-ACT-GGA-TGC-AAT-GTT-AAA-G-3'$$
.

(Asterisks show the location of mismatches and the underlined sequence shows the position of the altered Sau3A site.) The B. amyloliquefaciens subtilisin gene on a 1.5 kb EcoRI-BAMHI fragment from pS4.5 was cloned into M13mp11 and single stranded DNA was isolated. This template (M13mp11SUBT) was double primed with the 5' phosphorylated M13 universal sequencing primer and the mutagenesis primer. Adelman, et al. (1983) DNA 2, 183-193. The heteroduplex was transfected into competent JM101 cells and plaques were probed for the mutant sequence (Zoller, M.J., et al. (1982) Nucleic Acid Res. 10, 6487-6500; Wallace, et al. (1981) Nucleic Acid Res. 9, 3647-3656) using a tetramethylammonium chloride hybridization protocol (Wood, et al. (1985) Proc. Natl. Acad. Sci. USA 82, 1585-1588). The Ser87 to Cys mutation was prepared in a similar fashion using a 5' phosphorylated primer having the sequence

(The asterisk indicates the position of the mismatch and the underlined sequence shows the position of a new Mstl site.) The C24 and C87 mutations were obtained at a frequency of one and two percent, respectively. Mutant sequences were confirmed by dideoxy sequencing in M13.

Mutagenesis of Tyr21/Thr22 to A21/C22 was carried out with a 5' phosphorylated oligonucleotide primer having the sequence

(The asterisks show mismatches to the wild type sequence and the underlined sequence shows the position of an altered <u>Sau3A</u> site.) Manipulations for heteroduplex synthesis were identical to those described for C24. Because direct cloning of the heteroduplex DNA fragment can yield increased frequencies of mutagenesis, the <u>EcoRI-BamHI</u> subtilisin fragment was purified and ligated into pBS42. <u>E. coli MM 294 cells were transformed with the ligation mixture and plasmid DNA was purified from isolated transformants. Plasmid DNA was screened for the loss of the <u>Sau3A</u> site at codon 23 that was eliminated by the mutagenesis primer. Two out of 16 plasmid preparations had lost the wild type <u>Sau3A</u> site. The mutant sequence was confirmed by dideoxy sequencing in M13.</u>

Double mutants, C22/C87 and C24/C87, were constructed by ligating fragments sharing a common Clais site that separated the single parent cystine codons. Specifically, the 500 bp EcoRI-Clai fragment containing the 5' portion of the subtilisin gene (including codons 22 and 24) was ligated with the 4.7 kb Clai-EcoRI fragment that contained the 3' portion of the subtilisin gene (including codon 87) plus pBS42 vector sequence. E. coli MM 294 was transformed with ligation mixtures and plasmid DNA was purified from individual transformants. Double-cysteine plasmid constructions were identified by restriction site markers originating from the parent cysteine mutants (i.e., C22 and C24, Sau3A minus; Cys87, Mstl plus). Plasmids from E. coli were transformed into B. subtilis BG2036. The thermal stability of these mutants as compared to wild type subtilisin are presented in Figure 30 and Tables XVIII and XVIII.

TABLE XVII

E	Effect of DTT on the Half-Time of Autolytic Inactivation of Wild-Type and Disulfide Mutants of Subtilisin*													
5	Enzyme	t,	-DTT/+DTT											
		-DDT	+DTT]										
		mi												
	Wild-type	95	85	1.1										
10	C22/C87	44	25	1.8										
	C24/C87	92	1.5											

(*) Purified enzymes were either treated or not treated with 25mM DTT and dialyzed with or without 10mM DTT in 2mM CaCl₂, 50mM Tris (pH 7.5) for 14 hr. at 4 °C. Enzyme concentrations were adjusted to 80µI aliquots were quenched on ice and assayed for residual activity. Half-times for autolytic inactivation were determined from semi-log plots of log₁₀ (residual activity) versus time. These plots were linear for over 90% of the inactivation.

TABLE XVIII

Effect of Mutations in Subtilisin on the Half-Time of Autolytic Inactivation at 58 ° C*									
Enzyme	tį								
	min								
Wild-type	120								
C22	22								
C24	120								
C87	104								
C22/C87	43								
C24/C87	115								

(*) Half-times for autolytic inactivation were determined for wild-type and mutant subtilisins as described in the legend to Table III. Unpurified and non-reduced enzymes were used directly from B. subtilis culture supernatants.

The disulfides introduced into subtilisin did not improve the autolytic stability of the mutant enzymes when compared to the wild-type enzyme. However, the disulfide bonds did provide a margin of autolytic stability when compared to their corresponding reduced double-cysteine enzyme. Inspection of a highly refined x-ray structure of wild-type B. amyloliquefaciens subtilisin reveals a hydrogen bond between Thr22 and Ser87. Because cysteine is a poor hydrogen donor or acceptor (Paul, I.C. (1974) in Chemistry of the -SH Group (Patai, S., ed.) pp. 111-149, Wiley Interscience, New York) weakening of 22/87 hydrogen bond may explain why the C22 and C87 single-cysteine mutant proteins are less autolytically stable than either C24 or wild-type (Table XVIII). The fact that C22 is less autolytically stable than C87 may be the result of the Tyr21A mutation (Table XVIII). Indeed, construction and analysis of Tyr21/C22 shows the mutant protein has an autolytic stability closer to that of C87. In summary, the C22 and C87 of single-cysteine mutations destabilize the protein toward autolysis, and disulfide bond formation increases the stability to a level less than or equal to that of wild-type enzyme.

EXAMPLE 12

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Multiple Mutants Containing Substitutions at Position 222 and Position 166 or 169

Double mutants 166/222 and 169/222 were prepared by ligating together (1) the 2.3kb Acall fragment from pS4.5 which contains the 5' portion of the subtilisin gene and vector sequences, (2) the 200bp Avall fragment which contains the relevant 166 or 169 mutations from the respective 166 or 169 plasmids, and (3) the 2.2kb Avall fragment which contains the relevant 222 mutation 3' and of the subtilisin genes and vector

sequence from the respective p222 plasmid.

Although mutations at position 222 improve oxidation stability they also tend to increase the Km. An example is shown in Table XIX. In this case the A222 mutation was combined with the K166 mutation to give an enzyme with kcat and Km intermediate between the two parent enzymes.

TABLE XIX

	kçat	Km								
WT	50	1.4x10 ⁻⁴								
A222	42	9.9x10 ⁻⁴								
K166	21	3.7x10 ⁻⁵								
K166/A222	29	2.0x10 ⁻⁴								
substrate sAAPFpNa										

EXAMPLE 13

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Multiple Mutants Containing Substitutions at Positions 50, 156, 166, 217 and Combinations Thereof

The double mutant S156/A169 was prepared by ligation of two fragments, each containing one of the relevant mutations. The plasmid pS156 was cut with Xmall and treated with S1 nuclease to create a blunt end at codon 167. After removal of the nuclease by phenol/chloroform extraction and ethanol precipitation, the DNA was digested with BamHI and the approximately 4kb fragment containing the vector plus the 5' portion of the subtilisin gene through codon 167 was purified.

The pA169 plasmid was digested with KpnI and treated with DNA polymerase Klenow fragment plus 50 μ M dNTPs to create a blunt end codon at codon 168. The Klenow was removed by phenol/chloroform extraction and ethanol precipitation. The DNA was digested with BamHI and the 590bp fragment including codon 168 through the carboxy terminus of the subtilisin gene was isolated. The two fragments were then ligated to give S156/A169.

Triple and quadruple mutants were prepared by ligating together (1) the 220bp Pvull/HaeII fragment containing the relevant 156, 166 and/or 169 mutations from the respective p156, p166 and/or p169 double of single mutant plasmid, (2) the 550bp HaeII/BamHI fragment containing the relevant 217 mutant from the respective p217 plasmid, and (3) the 3.9kb Pvull/BamHI fragment containing the F50 mutation and vector sequences.

The multiple mutant F50/S156/A169/L217, as well as <u>B. amyloliquefaciens</u> subtilisin, <u>B. lichenformis</u> subtilisin and the single mutant L217 were analyzed with the above synthetic polypeptides where the P-1 amino acid in the substrate was Lys, His, Ala, Gln, Tyr, Phe, Met and Leu. These results are shown in Figures 26 and 27.

These results show that the F50/S156/A169/L217 mutant has substrate specificity similar to that of the B. <u>licheniformis</u> enzyme and differs dramatically from the wild type enzyme. Although only data for the L217 mutant are shown, none of the single mutants (e.g., F50, S156 or A169) showed this effect. Although B. <u>licheniformis</u> differs in 88 residue positions from B. <u>amyloliquefaciens</u>, the combination of only these four mutations accounts for most of the differences in substrate specificity between the two enzymes.

EXAMPLE 14

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Subtilisin Mutants Having Altered Alkaline Stability

A random mutagenesis technique was used to generate single and multiple mutations within the <u>B. amyloliquefaciens</u> subtilisin gene. Such mutants were screened for altered alkaline stability. Clones having increased (positive) alkaline stability and decreased (negative) alkaline stability were isolated and sequenced to identify the mutations within the subtilisin gene. Among the positive clones, the mutants V107 and R213 were identified. These single mutants were subsequently combined to produce the mutant V107/R213.

One of the negative clones (V50) from the random mutagenesis experiments resulted in a marked decrease in alkaline stability. Another mutant (P50) was analyzed for alkaline stability to determine the effect

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of a different substitution at position 50. The F50 mutant was found to have a greater alkaline stability than wild type subtilisin and when combined with the double mutant V107/R213 resulted in a mutant having an alkaline stability which reflected the aggregate of the alkaline stabilities for each of the individual mutants.

The single mutant R204 and double mutant C204/R213 were identified by alkaline screening after random cassette mutagenesis over the region from position 197 to 228. The C204/R213 mutant was thereafter modified to produce mutants containing the individual mutations C204 and R213 to determine the contribution of each of the individual mutations. Cassette mutagenesis using pooled oligonucleotides to substitute all amino acids at position 204, was utilized to determine which substitution at position 204 would maximize the increase in alkaline stability. The mutation from Lys213 to Arg was maintained constant for each of these substitutions at position 204.

A. Construction of pB0180, an E. coli-B. subtilis Shuttle Plasmid

The 2.9 kb EcoRI-BamHI fragment from pBR327 (Covarrubias, L., et al. (1981) Gene 13, 25-35) was ligated to the 3.7kb EcoRI-BamHI fragment of pBD64 (Gryczan, T., et al. (1980) J. Bacteriol., 141, 246-253) to give the recombinant plasmid pB0153. The unique EcoRI recognition sequence in pBD64 was eliminated by digestion with EcoRI followed by treatment with Klenow and deoxynucleotide triphosphates (Maniatis, T., et al. (eds.) (1982) in Molecular Cloning, A Laboratory Manual, Cold spring Harbor Laboratory, Cold Spring Harbor, N.Y.). Blunt end ligation and transformation yielded pB0154. The unique Aval recognition sequence in pBO154 was eliminated in a similar manner to yield pBO171, pB0171 was digested with BamHI and Pvull and treated with Klenow and deoxynucleotide triphosphates to create blunt ends. The 6.4 kb fragment was purified, ligated and transformed into LE392 cells (Enquest, L.W., et al. (1977) J. Mol. Biol. 111, 97-120), to yield pB0172 which retains the unique BamHI site. To facilitate subcloning of subtilisin mutants, a unique and silent KpnI site starting at codon 166 was introduced into the subtilisin gene from pS4.5 (Wells, J.A., et al. (1983) Nucleic Acids Res., 11, 7911-7925) by site-directed mutagenesis. The Kpnl+ plasmid was digested with EcoRI and treated with Klenow and deoxynucleotide triphosphates to create a blunt end. The Klenow was inactivated by heating for 20 min at 68 °C, and the DNA was digested with BamHI. The 1.5 kb blunt EcoRI-BamHI fragment containing the entire subtilisin was ligated with the 5.8 kb Nrul-BamHI from pB0172 to yield pB0180. The ligation of the blunt Nrul end to the blunt EcoRI end recreated an EcoRI site. Proceeding clockwise around pB0180 from the EcoRI site at the 5' end of the subtilisin gene is the unique BamHI site at the 3' end of the subtilisin gene, the chloramphenicol and neomycin resistance genes and UB110 gram positive replication origin derived from pBD64, the ampicillin resistance gene and gram negative replication origin derived from pBR327.

5 B. Construction of Random Mutagenesis Library

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The 1.5 kb EcoRI-BamHI fragment containing the B. amyloliquefaciens subtilisin gene (Wells et al., 1983) from pB0180 was cloned into M13mp11 to give M13mp11 SUBT essentially as previously described (Wells, J.A., et al. (1986) J. Biol. Chem., 261,6564-6570). Deoxyuridine containing template DNA was prepared according to Kunkel (Kunkel, T.A. (1985) Proc. Natl. Acad. Sci. USA, 82 488-492). Uridine containing template DNA (Kunkel, 1985) was purified by CsCI density gradients (Maniatis, T. et al. (eds.) (1982) in Molecular Cloning, A Laboratory Manual, Cold Spring Harbor Laboratory, Cold Spring Harbor, N.Y.). A primer (Aval⁻) having the sequence

5 GAAAAAAGACCCTAGCGTCGCTTA

ending at codon -11, was used to alter the unique <u>Aval</u> recognition sequence within the subtilisin gene. (The asterisk denotes the mismatches from the wild-type sequence and underlined is the altered Aval site.)

The 5' phosphorylated Aval primer (~320 pmol) and ~40 pmol (~120µg) of uridine containing M13mp11 SUBT template in 1.88 ml of 53 mM NaCl, 7.4 mM MgCl2 and 7.4 mM Tris.HCl (pH 7.5) were annealed by heating to 90 °C for 2 min. and cooling 15 min at 24 °C (Fig. 31). Primer extension at 24 °C was initiated by addition of 100µL containing 1 mM in all four deoxynucleotide triphosphates, and 20µl Klenow fragment (5 units/l). The extension reaction was stopped every 15 seconds over ten min by addition of 10µl 0.25 M EDTA (pH 8) to 50µl aliquots of the reaction mixture. Samples were pooled, phenol chlorophorm extracted and DNA was precipitated twice by addition of 2.5 vol 100% ethanol, and washed twice with 70% ethanol.

The pellet was dried, and redissolved in 0.4 ml 1 mM EDTA, 10 mM Tris (pH 8).

Misincorporation of α -thiodeoxynucleotides onto the 3' ends of the pool of randomly terminated template was carried out by incubating four 0.2 ml solutions each containing one-fourth of the randomly terminated template mixture (~20 μ g), 0.25 mM of a given α -thiodeoxynucleotide triphosphate, 100 units AMV polymerase, 50 mM KCL, 10 mM MgCl₂, 0.4 mM dithiothreitol, and 50 mM Tris (pH 8.3) (Champoux, J.J. (1984) Genetics, 2, 454-464). After incubation at 37 °C for 90 minutes, misincorporation reactions were sealed by incubation for five minutes at 37 °C with 50 mM all four deoxynucleotide triphosphates (pH 8), and 50 units AMV polymerase. Reactions were stopped by addition of 25 mM EDTA (final), and heated at 68 °C for ten min to inactivate AMV polymerase. After ethanol precipitation and resuspension, synthesis of closed circular heteroduplexes was carried out for two days at 14 °C under the same conditions used for the timed extension reactions above, except the reactions also contained 1000 units T4 DNA ligase, 0.5 mM ATP and 1 mM β -mercaptoethanol. Simultaneous restriction of each heteroduplex pool with Kpnl, BamHl, and EcoRI confirmed that the extension reactions were nearly quantitative. Heteroduplex DNA in each reaction mixture was methylated by incubation with 80 μ M S-adenosylmethionine and 150 units dam methylase for 1 hour at 37 °C. Methylation reactions were stopped by heating at 68 °C for 15 min.

One-half of each of the four methylated heteroduplex reactions were transformed into 2.5 ml competent E. coli JM101 (Messing, J. (1979) Recombinant DNA Tech. Bull., 2, 43-48). The number of independent transformants from each of the four transformations ranged from 0.4-2.0 x 10^5 . After growing out phage pools, RF DNA from each of the four transformations was isolated and purified by centrifugation through CsCl density gradients. Approximately $2\mu g$ of RF DNA from each of the four pools was digested with EcoRI, BamHI and Aval. The 1.5 kb EcoRI-BamHI fragment (i.e., Aval resistant) was purified on low gel temperature agarose and ligated into the 5.5 kb EcoRI-BamHI vector fragment of pB0180. The total number of independent transformants from each α -thiodeoxynucleotide misincorporation plasmid library ranged from 1.2-2.4 x 10^4 . The pool of plasmids from each of the four transformations was grown out in 200 ml LB media containing 12.5 $\mu g/ml$ cmp and plasmid DNA was purified by centrifugation through CsCl density gradients.

C. Expression and Screening of Subtilisin Point Mutants

Plasmid DNA from each of the four misincorporation pools was transformed (Anagnostopoulos, C., et al. (1967), J. Bacteriol., 81, 741-746) into BG2036. For each transformation, 5µg of DNA produced approximately 2.5 x 105 independent BG2036 transformants, and liquid culture aliquots from the four libraries were stored in 10% glycerol at 70 °C. Thawed aliquots of frozen cultures were plated on LB/5µg/ml cmp/1.6% skim milk plates (Wells, J.A., et al. (1983) Nucleic Acids Res., 11, 7911-7925), and fresh colonies were arrayed onto 96-well microtiter plates containing 150 I per well LB media plus 12.5µg/ml cmp. After 1 h at room temperature, a replica was stamped (using a matched 96 prong stamp) onto a 132 mm BA 85 nitrocellulose filter (Schleicher and Scheull) which was layered on a 140 mm diameter LB/cmp/skim milk plate. Cells were grown about 16 h at 30 °C until halos of proteolysis were roughly 5-7 mm in diameter and filters were transferred directly to a freshly prepared agar plate at 37 °C containing only 1.6% skim milk and 50 mM sodium phosphate pH 11.5. Filters were incubated on plates for 3-6 h at 37 °C to produce halos of about 5 mm for wild-type subtilisin and were discarded. The plates were stained for 10 min at 24°C with Coomassie blue solution (0.25% Coomassie blue (R-250) 25% ethanol) and destained with 25% ethanol, 10% acetic acid for 20 min. Zones of proteolysis appeared as blue halos on a white background on the underside of the plate and were compared to the original growth plate that was similarly stained and destained as a control. Clones were considered positive that produced proportionately larger zones of proteolysis on the high pH plates relative to the original growth plate. Negative clones gave smaller halos under alkaline conditions. Positive and negative clones were restreaked to colony purify and screened again in triplicate to confirm alkaline pH results.

D. Identification and Analysis of Mutant Subtilisins

Plasmid DNA from 5 ml overnight cultures of more alkaline active B.subtilis clones was prepared according to Birnboim and Doly (Birnboim, H.C., et al. (1979) Nucleic Acid Res. 7, 1513) except that incubation with 2 mg/ml lysozyme proceeded for 5 min at 37 °C to ensure cell lysis and an additional phenol/CHCl₃ extraction was employed to remove contaminants. The 1.5 kb EcoRl-BamHI fragment containing the subtilisin gene was ligated into M13mp11 and template DNA was prepared for DNA sequencing (Messing, J., et al. (1982) Gene, 19 269-276). Three DNA sequence primers ending at codon 26, +95, and +155 were synthesized to match the subtilisin coding sequence. For preliminary sequence

identification a single track of DNA sequence, corresponding to the dNTPaS misincorporation library from which the mutant came, was applied over the entire mature protein coding sequence (i.e., a single dideoxyguanosine sequence track was applied to identify a mutant from the dGTPas library). A complete four track of DNA sequence was performed 200 bp over the site of mutagenesis to confirm and identify the mutant sequence (Sanger, F., et al., (1980) J. Mol. Biol., 143, 161-178). Confirmed positive and negative bacilli clones were cultured in LB media containing 12.5µg/mL cmp and purified from culture supernatants as previously described (Estell, D.A., et al. (1985) J. Biol. Chem., 260, 6518-6521). Enzymes were greater than 98% pure as analyzed by SDS-polyacrylamide gel electrophoresis (Laemmli, U.K. (1970), Nature, 227, 680-685), and protein concentrations were calculated from the absorbance at 280 nm,

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$$\varepsilon_{280}^{0.1\%} = 1.17$$

(Maturbara, H., et al. (1965), J. Biol. Chem, 240, 1125-1130).

Enzyme activity was measured with 200μg/mL succinyl-L-AlaL-AlaL-ProL-Phep-nitroanilide (Sigma) in 0.1M Tris pH 8.6 or 0.1 M CAPS pH 10.8 at 25 °C. Specific activity (μ moles product/min-mg) was calculated from the change in absorbance at 410 nm from production of p-nitroaniline with time per mg of enzyme (E410 = 8,480 M-lcm-l; Del Mar, E.G., et al. (1979), Anal. Biochem., 99, 316-320). Alkaline autolytic stability studies were performed on purified enzymes (200μg/mL) in 0.1 M potassium phosphate (pH 12.0) at 37 °C. At various times aliquots were assayed for residual enzyme activity (Wells, J.A., et al. (1986) J. Biol. Chem., 261, 6564-6570).

E. Results

1. Optimization and analysis of mutagenesis frequency

A set of primer-template molecules that were randomly 3'-terminated over the subtilisin gene (Fig. 31) was produced by variable extension from a fixed 5'-primer (The primer mutated a unique Aval site at codon 11 in the subtilisin gene). This was achieved by stopping polymerase reactions with EDTA after various times of extension. The extent and distribution of duplex formation over the 1 kb subtilisin gene fragment was assessed by multiple restriction digestion (not shown). For example, production of new Hinfl fragments identified when polymerase extension had proceeded past Ile110, Leu233, and Asp259 in the subtilisin gene.

Misincorporation of each dNTPαs at randomly terminated 3' ends by AMV reverse transcriptase (Zakour, R.A., et al. (1982), Nature, 295, 708-710; Zakour, R.A., et al. (1984), Nucleic Acids Res., 12, 6615-6628) used conditions previously described (Champoux, J.J., (1984), Genetics, 2, 454-464). The efficiency of each misincorporation reaction was estimated to be greater than 80% by the addition of each dNTPαs to the Aval restriction primer, and analysis by polyacrylamide gel electrophoresis. Misincorporations were sealed by polymerization with all four dNTP's and closed circular DNA was produced by reaction with DNA ligase.

Several manipulations were employed to maximize the yield of the mutant sequences in the heteroduplex. These included the use of a deoxyuridine containing template (Kunkel, T.A. (1985), Proc. Natl. Acad. Sci. USA, 82 488-492; Pukkila, P.J. et al. (1983), Genetics, 104, 571-582), in vitro methylation of the mutagenic strand (Kramer, W. et al. (1982) Nucleic Acids Res., 10 6475-6485), and the use of Aval restriction-selection against the wild-type template strand which contained a unique Aval site. The separate contribution of each of these enrichment procedures to the final mutagenesis frequency was not determined, except that prior to Aval restriction-selection roughly one-third of the segregated clones in each of the four pools still retained a wild-type Aval site within the subtilisin gene. After Aval restriction-selection greater than 98% of the plasmids lacked the wild-type Aval site.

The 1.5 kb EcoRI-BamHI subtilisin gene fragment that was resistant to Aval restriction digestion, from each of the four CsCI purified M13 RF pools was isolated on low melting agarose. The fragment was ligated in situ from the agarose with a similarly cut E. coli-B. subtilis shuttle vector, pB0180, and transformed directly into E coli LE392. Such direct ligation and transformation of DNA isolated from agarose avoided loses and allowed large numbers of recombinants to be obtained (>100,000 per µg equivalent of input M13 pool).

The frequency of mutagenesis for each of the four dNTPas misincorporation reactions was estimated from the frequency that unique restriction sites were eliminated (Table XX). The unique restriction sites

chosen for this analysis, <u>Clal</u>, <u>Pvull</u>, and <u>Kpnl</u>, were distributed over the subtilisin gene starting at codons 35, 104, and 166, respectively. As a control, the mutagenesis frequency was determined at the <u>Pstl</u> site located in the <u>B lactamase</u> gene which was outside the window of mutagenesis. Because the absolute mutagenesis frequency was close to the percentage of undigested plasmid DNA, two rounds of restriction-selection were necessary to reduce the background of surviving uncut wild-type plasmid DNA below the mutant plasmid (Table XX). The background of surviving plasmid from wild-type DNA probably represents the sum total of spontaneous mutations, uncut wild-type plasmid, plus the efficiency with which linear DNA can transform <u>E. coli</u>. Subtracting the frequency for unmutagenized DNA (background) from the frequency for mutant DNA, and normalizing for the window of mutagenesis sampled by a given restriction analysis (4-6 bp) provides an estimate of the mutagenesis efficiency over the entire coding sequence (~1000 bp).

TABLE XX

5	a-thiol dNTP misincor- porated (b)	Restriction Site Selection	% resi	stant o 2nd round	Total	% resistant clones over Background ^d	mutants per 1000bp		
	None	PstI	0.32	0.7	0.002	0	-		
10	G	PstI	0.33	1.0	0.003	0.001	0.2		
	Ŧ	PstI	0.32	<0.5	<0.002	0	0		
	С	PstI	0.43	3.0	0.013	0.011	3		
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	None	<u>Cla</u> I	0.28	5	0.014	0	-		
	G	<u>Cla</u> I	2.26	85	1.92	1.91	380		
	T	<u>Cla</u> I	0.48	31	0.15	0.14	35		
20	С	ClaI	0.55	15	0.08	0.066	17		
	None	PvuII	0.08	29	0.023	0	_		
25	G	PvuII	0.41	90	0.37	0.35	88		
25	T	PvuII	0.10	67	0.067	0.044	9		
	c	<u>Pvu</u> II	0.76	53	0.40	0.38	95		
30	None	KpnI	0.41	3	0.012	0	-		
	G	KpnI	0.98	3 5	0.34	0.33	83		
	T	KpnI	0.36	15	0.054	0.042	8		
	C .	Kpnl	1.47	26	0.38	0.37	93		
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⁽a) Mutagenesis frequency is estimated from the frequency for obtaining mutations that alter unique restriction sites within the mutagenized subtilisin gene (i.e., ClaI, PvuII, or KpnI) compared to mutation frequencies of the PstI site, that is outside the window of mutagenesis.

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⁽b) Plasmid DNA was from wild-type (none) or mutagenized by dNTPas misincorporation as described.

⁽c) Percentage of resistant clones was calculated from the fraction of clones obtained after three fold or greater over-digestion of the plasmid with the indicated restriction enzyme compared to a

non-digested control. Restriction-resistant plasmid DNA from the first round was subjected to a second round of restriction-selection. The total represents the product of the fractions of resistant clones obtained from both rounds of selection and gives percentage of restriction-site mutant clones in the original starting pool. Frequencies were derived from counting at least 20 colonies and usually greater than 100.

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- (d) Percent resistant clones was calculated by subtracting the percentage of restriction-resistant clones obtained for wild-type DNA (i.e., none) from that obtained for mutant DNA.
- (e) This extrapolates from the frequency of mutation over each restriction site to the entire subtilisin gene (~1 kb). This has been normalized to the number of possible bases (4-6 bp) within each restriction site that can be mutagenized by a given misincorporation event.

From this analysis, the average percentage of subtilisin genes containing mutations that result from dGTPas, dCTPas, or dTTPas misincorporation was estimated to be 90, 70, and 20 percent, respectively. These high mutagenesis frequencies were generally quite variable depending upon the dNTPas and misincorporation efficiencies at this site. Misincorporation efficiency has been reported to be both dependent on the kind of mismatch, and the context of primer (Champoux, J.J., (1984); Skinner, J.A., et al. (1986) Nucleic Acids Res., 14, 6945-6964). Biased misincorporation efficiency of dGTPas and dCTPas over dTTPas has been previously observed (Shortle, D., et al. (1985), Genetics, 110, 539-555). Unlike the dGTPas, dCTPas, and dTTPas libraries the efficiency of mutagenesis for the dATPas misincorporation library could not be accurately assessed because 90% of the restriction-resistant plasmids analyzed simply lacked the subtilisin gene insert. This problem probably arose from self-ligation of the vector when the dATPas mutagenized subtilisin gene was subcloned from M13 into pB0180. Correcting for the vector background, we estimate the mutagenesis frequency around 20 percent in the dATPas misincorporation library. In a separate experiment (not shown), the mutagenesis efficiencies for dGTPas and dTTPas misincorporation were estimated to be around 50 and 30 percent, respectively, based on the frequency of reversion of an inactivating mutation at codon 169.

The location and identity of each mutation was determined by a single track of DNA sequencing corresponding to the misincorporated athiodeoxynucleotide over the entire gene followed by a complete four track of DNA sequencing focused over the site of mutation. Of 14 mutants identified, the distribution was similar to that reported by Shortle and Lin (1985) except we did not observe nucleotide insertion or deletion mutations. The proportion of AG mutations was highest in the G misincorporation library, and some unexpected point mutations appeared in the dTTPas and dCTPas libraries.

2. Screening and Identification of Alkaline Stability Mutants of Subtilisin

It is possible to screen colonies producing subtilisin by halos of casein digestion (Wells, J.A. et al. (1983) Nucleic Acids Res., 11, 7911-7925). However, two problems were posed by screening colonies under high alkaline conditions (>pH 11). First, B. subtilis will not grow at high pH, and we have been unable to transform an alkylophilic strain of bacillus. This problem was overcome by adopting a replica plating strategy in which colonies were grown on filters at neutral pH to produce subtilisin and filters subsequently transferred to casein plates at pH 11.5 to assay subtilisin activity. However, at pH 11.5 the casein micells no longer formed a turbid background and thus prevented a clear observation of proteolysis halos. The problem was overcome by briefly staining the plate with Coomassie blue to amplify proteolysis zones and acidifying the plates to develop casein micell turbidity. By comparison of the halo size produced on the reference growth plate (pH 7) to the high pH plate (pH 11.5), it was possible to identify mutant subtilisins that had increased (positives) or decreased (negatives) stability under alkaline conditions.

Roughly 1000 colonies were screened from each of the four misincorporation libraries. The percentage of colonies showing a differential loss of activity at pH 11.5 versus pH 7 represented 1.4, 1.8, 1.4, and 0.6% of the total colonies screened from the thiol dGTPas, dATPas, dTTPas, and dCTPas libraries, respectively. Several of these negative clones were sequenced and all were found to contain a single base change as expected from the misincorporation library from which they came. Negative mutants included A36, E170 and V50. Two positive mutants were identified as V107 and R213. The ratio of negatives to positives was roughly 50:1.

3. Stability and Activity of Subtilisin Mutants at Alkaline pH

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Subtilisin mutants were purified and their autolytic stabilities were measured by the time course of inactivation at pH 12.0 (Figs. 32 and 33). Positive mutants identified from the screen (i.e., V107 and R213) were more resistant to alkaline induced autolytic inactivation compared to wild-type; negative mutants (i.e., E170 and V50) were less resistant. We had advantageously produced another mutant at position 50 (F50) by site-directed mutagenesis. This mutant was more stable than wild-type enzyme to alkaline autolytic inactivation (Fig. 33) At the termination of the autolysis study, SDS-PAGE analysis confirmed that each subtilisin variant had autolyzed to an extent consistent with the remaining enzyme activity.

The stabilizing effects of V107, R213, and F50 are cumulative. See Table XXI. The double mutant, V107/R213 (made by subcloning the 920 bp EcoRI-KpnI fragment of pB0180V107 into the 6.6 kb EcoRI-KpnI fragment of pB0180R213), is more stable than either single mutant. The triple mutant, F50/V107/R213 (made by subcloning the 735 bp EcoRI-Pvull fragment of pF50 (Example 2) into the 6.8 kb EcoRI-Pvull fragment of pB0180/V107, is more stable than the double mutant V107/R213 or F50. The inactivation curves show a biphasic character that becomes more pronounced the more stable the mutant analyzed. This may result from some destablizing chemical modification(s) (eg., deamidation) during the autolysis study and/or reduced stabilization caused by complete digestion of larger autolysis peptides. These alkaline autolysis studies have been repeated on separately purified enzyme batches with essentially the same results. Rates of autolysis should depend both on the conformational stability as well as the specific activity of the subtilisin variant (Wells, J.A., et al. (1986), J. Biol. Chem., 261, 6564-6570). It was therefore possible that the decreases in autolytic inactivation rates may result from decreases in specific activity of the more stable mutant under alkaline conditions. In general the opposite appears to be the case. The more stable mutants, if anything, have a relatively higher specific activity than wild-type under alkaline conditions and the less stable mutants have a relatively lower specific activity. These subtle effects on specific activity for V107/R213 and F50/V107/R213 are cumulative at both pH 8.6 and 10.8. The changes in specific activity may reflect slight differences in substrate specificity, however, it is noteworthy that only positions 170 and 107 are within 6A of a bound model substrate (Robertus, J.D., et al. (1972), Biochemistry 11, 2438-2449).

TABLE XXI

40	Relationship between relative specific acitivity at pH 8.6 or 10.8 and alkaline autolytic stability													
Ĭ	Enzyme	Relative sp	ecific activity	Alkaline autolysis half-time (min)b										
		pH 8.6	pH 10.8											
Γ	Wild-type	100±1	100±3	86										
45	Q170	46±1	28±2	13										
	V107	126±3	99±5	102										
	R213	97±1	102±1	115										
	V107/R213	116±2	106±3	130										
	V50	66±4	61±1	58										
50	F50	123±3	157±7	131										
	F50/V107/R213	126±2	152±3	168										

⁽a) Relative specific activity was the average from triplicate activity determinations divided by the wild-type value at the same pH. The average specific activity of wild-type enzyme at pH 8.6 and 10.8 was 70µmoles/min-mg and 37µmoles/min-mg, respectively.

(b) Time to reach 50% activity was taken from Figs. 32 and 33.

F. Random Cassette Mutagenesis of Residues 197 through 228

Plasmid p Δ 222 (Wells, et al. (1985) Gene 34, 315-323) was digested with Pstl and BamHl and the 0.4 kb Pstl/BamHl fragment (fragment 1, see Fig. 34) purified from a polyacrylamide gel by electroelution.

The 1.5 kb <u>EcoRI/Bam</u>HI fragment from pS4.5 was cloned into M13mp9. Site directed mutagenesis was performed to create the A197 mutant and simultaneously insert a silent <u>SstI</u> site over codons 195-196. The mutant <u>EcoRI/BamHI</u> fragment was cloned back into pBS42. The pA197 plasmid was digested with <u>Bam</u>HI and SstI and the 5.3 kb BamHI/SstI fragment (fragment 2) was purified from low melting agarose.

Complimentary oligonucleotides were synthesized to span the region from Sstl (codons 195-196) to Pstl (codons 228-230). These oligodeoxynucleotides were designed to (1) restore codon 197 to the wild type, (2) re-create a silent Kpnl site present in p∆222 at codons 219-220, (3) create a silent Smal site over codons 210-211, and (4) eliminate the Pstl site over codons 228-230 (see Fig. 35). Oligodeoxynucleotides were synthesized with 2% contaminating nucleotides at each cycle of synthesis, e.g., dATP reagent was spiked with 2% dCTP, 2% dGTP, and 2% dTTP. For 97-mers, this 2% poisoning should give the following percentages of non-mutant, single mutants and double or higher mutants per strand with two or more misincorporations per complimentary strand: 14% non-mutant, 28% single mutant, and 57% with ≥2 mutations, according to the general formula

$$f = \frac{\mu^n}{n!} e^{-\mu}.$$

where μ is the average number of mutations and n is a number class of mutations and f is the fraction of the total having that number of mutations. Complimentary oligodeoxynucleotide pools were phosphorylated and annealed (fragment 3) and then ligated at 2-fold molar excess over fragments 1 and 2 in a three-way ligation.

E. coli MM294 was transformed with the ligation reaction, the transformation pool-grown up over night and the pooled plasmid DNA was isolated. This pool represented 3.4 x 10⁴ independent transformants. This plasmid pool was digested with Pstl and then used to retransform E. coli. A second plasmid pool was prepared and used to transform B. subtilis (BG2036). Approximately 40% of the BG2036 transformants actively expressed subtilisin as judged by halo-clearing on casein plates. Several of the non-expressing transformants were sequenced and found to have insertions or deletions in the synthetic cassettes. Expressing BG2036 mutants were arrayed in microtiter dishes with 150μI of LB/12.5μg/mL chloramphenicol (cmp) per well, incubated at 37 °C for 3-4 hours and then stamped in duplicate onto nitrocellulose filters laid on LB 1.5% skim milk/5μg/mL cmp plates and incubated overnight at 33 °C (until halos were approximately 4-8 mm in diameter). Filters were then lifted to stacks of filter paper saturated with 1 x Tide commercial grade detergent, 50 mM Na₂CO₃, pH 11.5 and incubated at 65 °C for 90 min. Overnight growth plates were Commassie stained and destained to establish basal levels of expression. After this treatment, filters were returned to pH7/skim milk/20μg/mL tetracycline plates and incubated at 37 °C for 4 hours to overnight.

Mutants identified by the high pH stability screen to be more alkaline stable were purified and analyzed for autolytic stability at high pH or high temperature. The double mutant C204/R213 was more stable than wild type at either high pH or high temperature (Table XXII).

This mutant was dissected into single mutant parents (C204 and R213) by cutting at the unique <u>Smal</u> restriction site (Fig. 35) and either ligating wild type sequence 3' to the <u>Smal</u> site to create the single <u>C204</u> mutant or ligating wild type sequence 5' to the <u>Smal</u> site to create the single R213 mutant. Of the two single parents, C204 was nearly as alkaline stable as the parent double mutant (C04/R213) and slightly more thermally stable. See Table XXII. The R213 mutant was only slightly more stable than wild type under both conditions (not shown).

Another mutant identified from the screen of the 197 to 228 random cassette mutagenesis was R204. This mutant was more stable than wild type at both high pH and high temperature but less stable than C204.

TABLE XXII

Stability of subtilisin variants

Purified enzymes $(200\mu g/mL)$ were incubated in 0.1M phosphate, pH 12 at 30°C for alkaline autolysis, or in 2mM CaCl₂, 50mM MOPS, pH 7.0 at 62°C for thermal autolysis. At various times samples were assayed for residual enzyme activity. Inactivations were roughly pseudo-first order, and t 1/2 gives the time it took to reach 50% of the starting activity in two separate experiments.

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		t : (alka: auto:		t 1/2 (thermal autolysis)				
25	Subtilisin variant	*1	Exp. #2	Exp. _#1_	Exp.			
	wild type	30	25	20	23			
30	F50/V107/R213	49	41	18	23			
	R204	35	32	24	27			
	C204	43	46	38	40			
35	C204/R213	50	52	32	36			
	L204/R213	32	30	20	21			

G. Random Mutagenesis at Codon 204

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Based on the above results, codon 204 was targeted for random mutagenesis. Mutagenic DNA cassettes (for codon at 204) all contained a fixed R213 mutation which was found to slightly augment the stability of the C204 mutant.

Plasmid DNA encoding the subtilisin mutant C204/R213 was digested with <u>Sstl</u> and <u>EcoRl</u> and a 1.0 kb EcoRl/Sstl fragment was isolated by electro-elution from polyacrylamide gel (fragment 1, see Fig. 35).

C204/R213 was also digested with Smal and EcoRI and the large 4.7 kb fragment, including vector sequences and the 3' portion of coding region, was isolated from low melting agarose (fragment 2, see Fig. 36).

Fragments 1 and 2 were combined in four separate three-way ligations with heterophosphorylated fragments 3 (see Figs. 36 and 37). This heterophosphorylation of synthetic duplexes should preferentially drive the phosphorylated strand into the plasmid ligation product. Four plasmid pools, corresponding to the four ligations, were restricted with <u>Smal</u> in order to linearize any single cut C204/R213 present from fragment 2 isolation, thus reducing the background of C204/R213. <u>E. coli</u> was then re-transformed with

<u>Smal</u>-restricted plasmid pools to yield a second set of plasmid pools which are essentially free of C204/R213 and any non-segregated heterduplex material.

These second enriched plasmid pools were then used to transform <u>B. subtilis</u> (BG2036) and the resulting four mutant pools were screened for clones expressing subtilisin resistant to high pH/temperature inactivation. Mutants found positive by such a screen were further characterized and identified by sequencing.

The mutant L204/R213 was found to be slightly more stable than the wild type subtilisin. See Table XXII.

Having described the preferred embodiments of the present invention, it will appear to those ordinarily skilled in the art that various modifications may be made to the disclosed embodiments, and that such modifications are intended to be within the scope of the present invention.

Claims

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- A subtilisin mutant derived by the substitution of at least one amino acid residue of a precursor subtilisin with a different amino acid, so that the subtilisin mutant has at least one property which is different from the same property of the precursor subtilisin, characterised by the substitution at one or more of Tyr21, Thr22, Ser24, Asp36, Ala45, Gly46, Ala48, Ser49, Met50, Asn77, Ser87, Lys94, Val95, Leu96, Ile107, Gly110, Met124, Lys170, Tyr171, Pro172, Asp197, Met199, Ser204, Lys213, His67, Leu135, Gly97, Ser101, Gly102, Glu103, Gly127, Gly128, Pro129, Tyr214, and Gly215 of Bacillus amyloliquefaciens subtilisin and equivalent amino acid residues in other precursor subtilisins.
- 2. A subtilisin mutant having an amino acid sequence derived from the amino acid sequence of a precursor subtilisin by the substitution of more than one amino acid residue of said amino acid sequence of said precursor subtilisin by a different amino acid, so that the subtilisin mutant has at least one property which is different from the same property of the precursor subtilisin, characterized by substitutions at more than one of Tyr21, Thr22, Ser24, Asp32, Ser33, Asp36, Ala45, Ala48, Ser49, Met50, Ser87, Lys94, Val95, Tyr104, Ile107, Gly110, Met124, Ala152, Asn155, Glu156, Gly166, Gly169, Lys170, Tyr171, Pro172, Phe189, Asp197, Met199, Ser204, Lys213, Tyr217, Ser221, Met222, His67, Leu135, Gly97, Ser101, Gly102, Glu103, Gly127, Gly128, Pro129, Tyr214, and Gly215 of Bacillus amyloliquefaciens subtilisin and equivalent amino acid residues in other precursor subtilisins, with the proviso that when substitution is made at any residue in the group Asp32, Ser33, Tyr104, Ala152, Asn155, Glu156 Gly166, Gly169, Phe189, Tyr217 and Met222 a substitution is also made at at least one specified position not of that group.
 - 3. The mutant of claim 2 wherein said combinations are selected from Thr22/Ser87, Ser24/Ser87, Ala45/Ala48, Ser49/Lys94, Ser49/Val95, Met50/Val95, Met50/Gly110, Met50/Met124, Met50/Met222, Met124/Met222, Tyr21/Thr22, Met50/Met124/Met222, Tyr21/Thr22/Ser87, Met50/Glu156/Gly166/Tyr217, Met50/Glu156/Tyr217, Ile170/Lys213, Ser204/Lys213, Met50/Ile107/Lys213 and Ser24/Met50/Ile107/Glu156/Gly166/Gly169/Ser204/Lys213/Gly215/Tyr217.
 - 4. A subtilisin mutant derived by the deletion of one or more amino acid residues in a precursor subtilisin equivalent to 161-164 in B. <u>amyloliquetaciens</u> subtilisin, said deletion being made alone or in combination with substitutions in the amino acid sequence of the precursor subtilisin, and producing at least one property which is different from the same property of the precursor subtilisin.
 - 5. A subtilisin mutant having altered substrate specificity when compared to a precursor subtilisin, the mutant being derived by the substitution of a different amino acid at the residue equivalent to Leu + 126 of B. <u>amyloliquefaciens</u> subtilisin, alone or in combination with other substitutions or deletions in the amino acid sequence of the precursor subtilisin.
 - 6. A subtilisin mutant having altered substrate specificity when compared to a precursor subtilisin, the mutant being derived by the substitution of a different amino acid at the residue equivalent to Asp + 99 in B. <u>amyloliquefaciens</u> subtilisin, alone or in combination with other substitutions or deletions in the amino acid sequence of the precursor subtilisin.
 - 7. A DNA sequence encoding the mutant of any one of the preceding claims.

- 8. An expression vector containing the mutant DNA sequence of claim 7.
- 9. A host cell transformed with the expression vector or claim 8.

5 Patentansprüche

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- 1. Subtilisinmutante, die durch Substitution zumindest eines Aminosäurerests eines Vorläufer-Subtilisins durch eine davon verschiedene Aminosäure hergeleitet ist, sodaß die Subtilisinmutante zumindest eine Eigenschaft aufweist, die sich von der gleichen Eigenschaft des Vorläufer-Subtilisins unterscheidet, gekennzeichnet durch die Substitution an einem oder mehreren von Tyr21, Thr22, Ser24, Asp36, Ala45, Gly46, Ala48, Ser49, Met50, Asn77, Ser87, Lys94, Val95, Leu96, Ile107, Gly110, Met124, Lys170, Tyr171, Pro172, Asp197, Met199, Ser204, Lys213, His67, Leu135, Gly97, Ser101, Gly102, Glu103, Gly127, Gly128, Pro129, Tyr214 und Gly215 von Bacillus amyloliquefaciens-Subtilisin und äquivalenten Aminosäureresten in anderen Vorläufer-Subtilisinen.
- 2. Subtilisinmutante mit einer Aminosäuresequenz, die aus der Aminosäuresequenz eines Vorläufer-Subtilisins durch Substitution mehr als eines Aminosäurerests der Aminosäuresequenz des Vorläufer-Subtilisins durch eine davon verschiedene Aminosäure hergeleitet ist, sodaß die Subtilisinmutante zumindest eine Eigenschaft auWeist, die sich von der gleichen Eigenschaft des Vorläufer-Subtilisins unterscheidet, gekennzeichnet durch Substitutionen an mehr als einem von Tyr21, Thr22, Ser24, Asp32, Ser33, Asp36, Ala45, Ala48, Ser49, Met50, Ser87, Lys94, Val95, Tyr104, Ile107, Gly110, Met124, Ala152, Asn155, Glu156, Gly166, Gly169, Lys170, Tyr171, Pro172, Phe189, Asp197, Met199, Ser204, Lys213, Tyr217, Ser221, Met222, His67, Leu135, Gly97, Ser101, Gly102, Glu103, Gly127, Gly128, Pro129, Tyr214 und Gly215 von Bacillus amyloliquefaciens-Subtilisin und äquivalenten Aminosäureresten in anderen Vorläufer-Subtilisinen, mit der Maßgabe, daß bei einer Substitution an irgendeinem Rest in der Gruppe Asp32, Ser33, Tyr104, Ala152, Asn155, Glu156, Gly166, Gly169, Phe189, Tyr217 und Met222 eine Substitution auch an zumindest einer bestimmten Position durchgeführt wird, die nicht dieser Gruppe angehört.
- Mutante nach Anspruch 2, worin die Kombinationen aus Thr22/Ser87, Ser24/Ser87, Ala45/Ala48, Ser49/Lys94, Ser49/Val95, Met50/Val95, Met50/Gly110, Met50/Met124, Met50/Met222, Met124/Met222, Tyr21/Thr22, Met50/Met124/Met222, Tyr21/Tyr22/Ser87, Met50/Glu156/Gly166/Tyr217, Met50/Glu156/Tyr217, Ile170/Lys213, Ser204/Lys213, Met50/Ile107/Lys213 und Ser24/Met50/Ile107/Glu156/Gly166/Gly169/Ser204/Lys213/Gly215/Tyr217 ausgewählt sind.
 - 4. Subtilisinmutante, die durch Löschung eines oder mehrerer Aminosäurereste in einem Vorläufer-Subtilisin, das 161-164 in B. amyloliquefaciens-Subtilisin äquivalent ist, hergeleitet ist, wobei die Löschung entweder alleine oder in Kombination mit Substitutionen in der Aminosäuresequenz des Vorläufer-Subtilisins erfolgt, und zumindest eine Eigenschaft ergibt, die sich von der gleichen Eigenschaft des Vorläufer-Subtilisins unterscheidet.
 - 6. Subtilisinmutante mit geänderter Substratspezifität im Vergleich zu einem Vorläufersubtilisin, wobei die Mutante durch Substitution einer unterschiedlichen Aminosäure am Rest, der Leu + 126 von B. amyloliquefaciens-Subtilisin äquivalent ist, alleine oder in Kombination mit anderen Substitutionen oder Löschungen in der Aminosäuresequenz des Vorläufer-Subtilisins hergeleitet ist.
 - 6. Subtilisinmutante mit geänderter Substratspezifität im Vergleich zu einem Vorläufersubtilisin, wobei die Mutante durch Substitution einer unterschiedlichen Aminosäure am Rest, der Asp +99 im B. amyloliquefaciens-Subtilisin äquivalent ist, alleine oder in Kombination mit anderen Substitutionen oder Löschungen in der Aminosäuresequenz des Vorläufer-Subtilisins hergeleitet ist.
 - 7. DNA-Sequenz, die für die Mutante nach einem der vorhergehenden Ansprüche kodiert.
 - Expressionsvektor, der die Mutanten-DNA-Sequenz von Anspruch 7 enthält.
 - 9. Wirtszelle, die mit dem Expressionsvektor von Anspruch 8 transformiert ist.

Revendications

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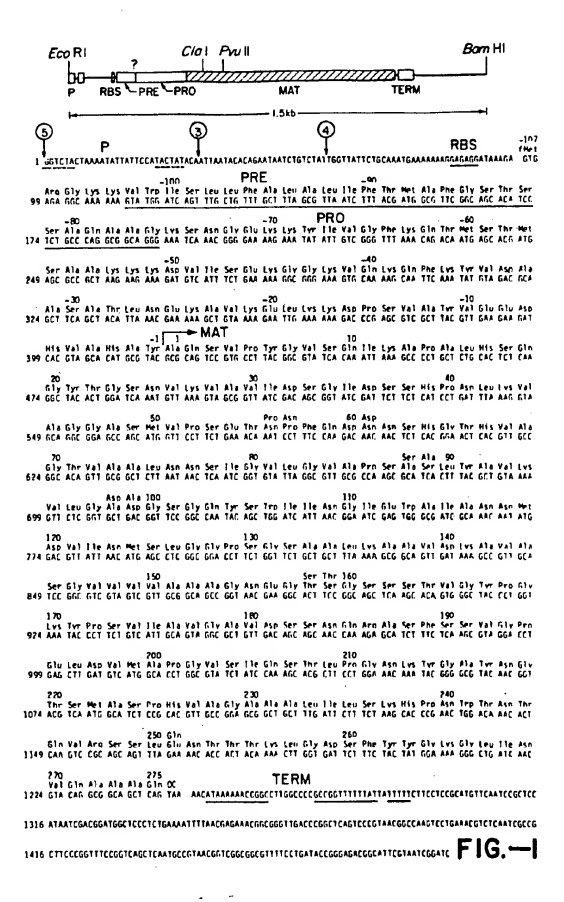
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- 1. Mutant de subtilisine dérivé par la substitution d'au moins un résidu d'acide aminé d'une subtilisine précurseur et par un acide aminé différent de manière que le mutant de subtilisine ait au moins une propriété qui est différente de la même propriété de la subtilisine précurseur, caractérisé par la substitution à un ou plusieurs de Tyr21, Thr22, Ser24, Asp36, Ala45, Gly46, Ala48, Ser49, Met50, Asn77, Ser87, Lys94, Val95, Leu96, Ile107, Gly110, Met124, Lys170, Tyr171, Pro172, Asp197, Met199, Ser204, Lys213, His67, Leu135, Gly97, Ser101, Gly102, Glu103, Gly127, Gly128, Pro129, Tyr214 et Gly215 de la subtilise de Bacillus amyloliquefaciens et les résidus d'acides aminés équivalents dans d'autres subtilisines précurseurs.
- 2. Mutant de subtilisine ayant une séquence d'acides aminés dérivée de la séquence d'acides aminés d'une subtilisine précurseur par la substitution de plus d'un résidu d'acide aminé de ladite séquence d'acides aminés de ladite subtilisine précurseur par un acide aminé différent de manière que le mutant de subtilisine ait au moins une propriété qui est différente de la même propriété de la subtilisine précurseur, caractérisé par des substitutions à plus d'un de Tyr21, Thr22, Ser24, Asp32, Ser33, Asp36, Ala45, Ala48, Ser49, Met50, Ser87, Lys94, Val95, Tyr104, Ile107, Gly110, Met124, Ala152, Asn155, Glu156, Gly166, Gly169, Lys170, Tyr171, Pro172, Phe189, Asp197, Met199, Ser204, Lys213, Tyr217, Ser221, Met222, His67, Leu135, Gly97, Ser101, Gly102, Glu103, Gly127, Gly128, Pro129, Tyr214 et Gly215 de la subtilisine de Bacillus amyloliquefaciens et des résidus d'acides aminés équivalents dans d'autres subtilisines précurseurs, à condition que quand la substitution est effectuée à tout résidu dans le groupe formé de Asp32, Ser33, Tyr104, Ala152, Asn155, Glu156, Gly166, Gly169, Phe189, Tyr217 et Met222, une substitution soit également effectuée en au moins une position spécifiée ne faisant pas partie de ce groupe.
- Mutant de la revendication 2 où lesdites associations sont choisies parmi Thr22/Ser87, Ser24/Ser87, Ala45/Ala48, Ser49/Lys94, Ser49/Val95, Met50/Val95, Met50/Gly110, Met50/Met124, Met50/Met222, Met124/Met222, Tyr21/Thr22, Met50/Met124/Met222, Tyr21/Thr22/ser87, Met50/Glu156/Gly166/Tyr217, Met50/Glu156/Tyr217, Ile170/Lys213, Ser204/Lys213, Met50/Ile107/Lys213 et Ser24/Met50/Ile107/Glu156/Gly166/Gly169/Ser204/Lys213/Gly215/Tyr217.
- 4. Mutant de subtilisine dérivé par la délétion d'un ou plusieurs résidus d'acides aminés dans une subtilisine précurseur équivalente à 161-164 dans la subtilisine de B. <u>amyloliquefaciens</u>, ladite délétion étant effectuée seule ou en association avec des substitutions dans la séquence d'acides aminés de la subtilisine précurseur et la production d'au moins une propriété qui est différente de la même propriété de la subtilisine précurseur.
- 5. Mutant de subtilisine ayant une spécificité modifiée du substrat en comparaison avec une subtilisine précurseur, le mutant étant dérivé par la substitution d'un acide aminé différent au résidu équivalent à Leu + 126 de la subtilisine de B. amyloliquefaciens, seule ou en association avec d'autres substitutions ou délétions dans la séquence d'acides aminés de la subtilisine précurseur.
- 6. Mutant de subtilisine ayant une spécificité modifiée de substrat en comparaison avec une subtilisine précurseur, le mutant étant dérivé par la substitution d'un acide aminé différent au résidu équivalent à Asp+99 dans la substitisine de <u>B. amyloliquefaciens</u>, seule ou en association avec d'autres substitutions ou délétions dans la séquence d'acides aminés de la subtilisine précurseur.
 - 7. Séquence d'ADN codant le mutant selon l'une quelconque des revendications précédentes.
- Vecteur d'expression contenant la séquence d'ADN du mutant de la revendication 7.
 - 9. Cellule hôte transformée par le vecteur d'expression de la revendication.8 .



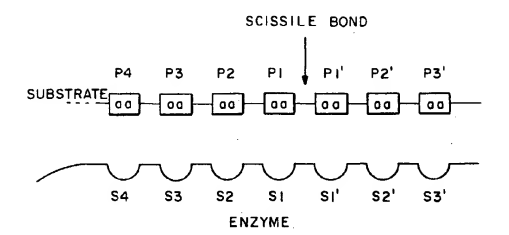


FIG. -2

FIG. - 3

F1G. -4

Honology of Bacillus protesses

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1.Bacillus amyloliquifaciens
2.Bacillus subtilis var.I158
3.Bacillus licheniformis (carlsbargensis)
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1 A A	0 0 0	5 5 T	v	P P	Y Y Y	6 6	V I	5 5 P	10 Q Q L	I	K	^ ^	P P D	6 6 K	r r	H	S S A	9	20 6 6
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61 N 6	N S N	5 5 6	H	6 6 6	T T	H	VVV	*	79 6 6 6	T T	V 1 V	6	6 6	L L L	N N D	N N N	5 5 1	I I T	80 6 6
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101 5 5	6 6 6	Q Q S	Y Y Y	5 5 5	6 n	1 1 1	I I V	N N S	110 6 6 6]]]	E E E	7 L	6 6	I I T	A 5 T	N N N	N N 6	M M M	120 D D

FIG. - 5A-1

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141 K K N	A A	V V Y	A S	5 S R	6 6	U I	VVV	VVV	150 V A	6 6	A A	A A A	6 6	N N N	E E S	6 5 6	T S N	\$ \$ \$	160 6 6
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241 U U L	† † 5	N N	T A S	0 0 0	U U U	R R R	S D N	S R R	250 L L L	E E S	N S S	T T T	T 6	T T T	K Y Y	i. L	6 6	D N S	260 S S
261 F F	Y Y Y	Y Y Y	6 6 6	K K K	6 6 6	L L L	I I	N N N	278 V V	0 0 E	♠ ♠	66	6 6	9 9 9					

FIG.-5A-2

ALIGNMENT OF 8.AMYLOLIQUIFACIENS SUBTILISIN AND THERMITASE 1.8. anyloliquifactons subtiliain 2.thermitees

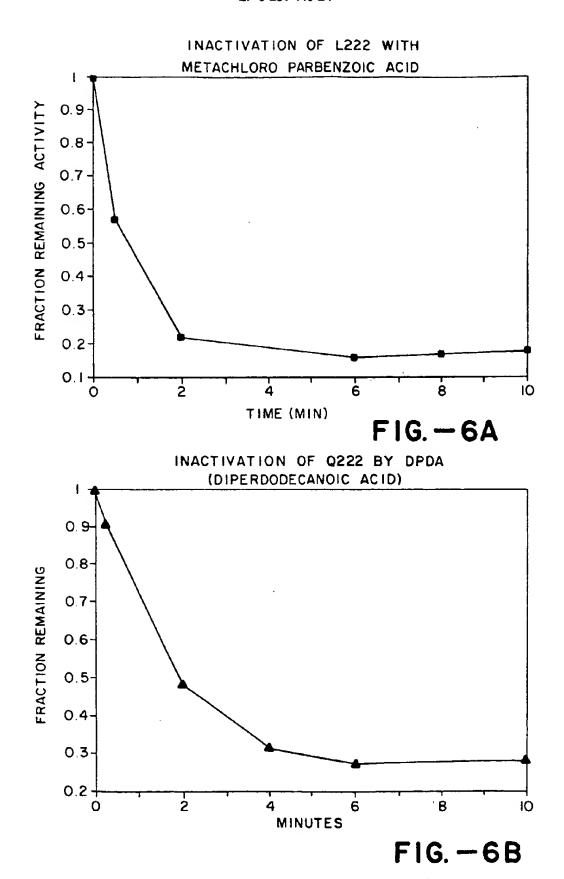
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									140										150
6	5 N	8	6	L	D K	A	A	V	D	K Y	A	U	A	5 K	6	Ų	V	V	150 V V

FIG. -5B-1

FIG. - 5B-2

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21	•	6	•	•	•	•	•	•	30	•	D	•	•	•	•	•		н	42
41	•		•		G		•	•	56	v	•	•	•	•	•	•	•	•	5 e
6 1	•	•	H	6	7	н	•		78 6		•	•	•	•	•	•	•	•	
8 1	•	6		•	•	•	•	•	•	•	•		•	Ų	L	•	•		188
101 S	•	•			•	•	•	•	1 1 8 6	•	•	•	•	•		•	•	•	128
121	•	•	•	•	L	6	•	•	130	•	•	•	•	•	•	•	•	•	148
141		•	•	•	ā	•	•		158	•	•	•	6	N	•	•	•	•	168
161	•	•	•	•	•	٧	P	•	170	•	•	•	•	•	•	v	•	•	186
181	•	•	•	•	•	•	8	F	190 5	•	•	•	•		•	•	•	•	208
281 P	6	•	•	•	•	•	•	•	216		•	•	•	•	•		•	6	728 T
		^	•	•	н	V	٨	c	. 536	•	•	•	•	•	•	•	•	•	248
241	•	•	•	·	•	R	•	•		•	•	•	•	٠	•	•	•	•	268
261	•	•	•		•		•	N	278										

FIG.-5C



73 02/12/2002, EAST Version: 1.03.0002

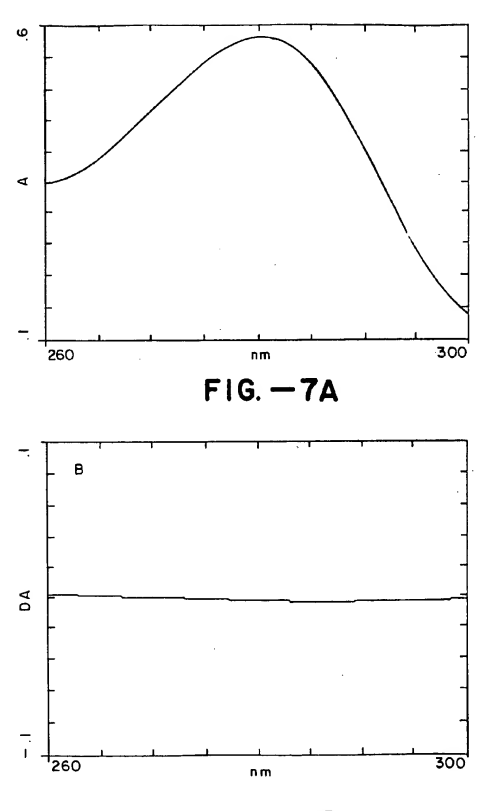


FIG. - 7B

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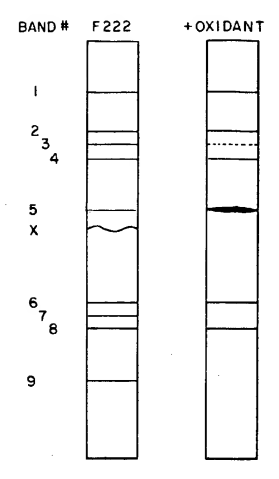


FIG. - 8

CNBr FRAGMENT MAP OF F222 MUTANT

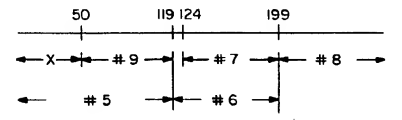


FIG. -9

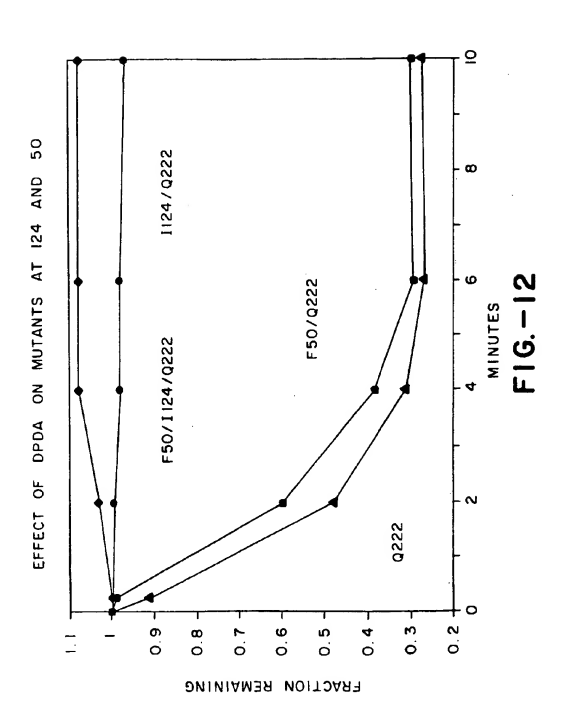
1. Codon number:	43 45
2. Wild type amino acid sequence:	Lys-Val-Ala-Gly-Gly-Ala-Ser-Met-Val-Pro-Ser
3. Wild type DNA sequence:	5'-AAG-GTA-GCA-GGC-GGA-GCC-AGC-ATG-GTT-CCT-TCT TTC-CAT-CGT-CCG-CCT-CGG-TCG-TAC-CAA-GGA-AGA-5'
4. p <u>∆</u> 50:	5'-AAG-GCC-TGC-ATG-GTA-CCT-TCT TTC-CGG-ACG-TAC-CAT-GGA-AGA-5'
5. pa50 cut with Stu I/Kpn I	5'-AAG-G TTC-Cp CAT-GGA-AGA-5'
6. Cut po50 ligated with cassettes:	* 5'-AAG-GTA-GCA-GGC-GGA-GCC-AGC-ATG-GTA-CCT-TCT TCC-CAT-CGT-CCG-CCT-CGG-TCG-TAC-CAT-GGA-AGA-5'
7. Mutagenesis primer for pΔ50:	*** 5'-ct-gat-tta-aag-gcc-tgc-atg-gta-cct-tct-ga
8. Mutants made:	V45, P45, V45/P48, E 46, E 48, V48, C 49, C 50, F 50

1. Codon number:	117 120 124 126 130
2. Wild type amino acid sequence:	INC8: Asn-Asn-Met-Asp-Val-Ile-Asn-Met-Ser-Leu-Gly-Gly-Pro-Ser
3. Wild type DNA sequence:	5'-AAC-AAT-ATG-GAC-GTT-ATT-AAC-ATG-AGC-CTC-GGC-GGA-CCT-TCT TTG-TTA-TAC-CTG-CAA-TAA-TTG-TAC-TCG-GAG-CCG-CCT-GGA-AGA-5'
4. p∆124:	* * * * * * * * * * * * * * * * * * *
5. p. b. 124 cut with Eco RV and Apa I	* pct-tct tig-tra-tra-trap ccg-gga-aga-5
6. Cut po124 ligated with cassettes:	* 5'-aac-aat-atg-gat-gtt-att-aac-atg-agc-ctc-ggc-ggc-cct-tct ttg-tta-tta-cta-caa-trg-tac-tcg-gag-ccg-gga-aga-5'
7. Mutagenesis primer for pΔ124::	5'-AAC-AAT-ATG-GAT-ATC-C-GGG-GGC-CCT-TCT-GGT-TC-3'

1. Codon number:

1124, L124 AND C126

8. Mutants made:



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3	Codon: Wild type amino acid sequence:	166 Thr Ser Giy Ser Ser Thr Val Gly Tyr Pro Gly
1:	Wild type DNA sequence:	5'-ACT TCC GGC AGC TCA AGC ACA GTG GGC TAC CCT GGT-3' 3'-TGA AGG CCG TCG AGT TCG TGT CAC CCG ATG GGA CCA-5'
2.	2. pal66 DNA sequence:	5'-ACT TCC 666 AGC TCA A C C CC6 6GT-3' 3'-TGA AGG CCC TCG AGT T G GGC CCA-5' SacI
ë.	3. pal66 cut with Sacl and Xmal: 5'-ACT TCC 666 A6C T	5'-ACT TCC 666 AGC T pcc6 66T-3' 3'-TGA AGG CCCp CA-5'
4.	Cut pal66 ligated with duplex DNA cassette pools:	5'-ACT TCC GGG AGC T <u>CA AGC ACA GTG NNN TAC</u> CCG GGT-3' 3'-TGA AGG CCC T <u>CG AGT TCG TGT CAC</u> NNN ATG GGC CCA-5'

MUTAGENESIS PRIMER 37 MER AA GGC ACT TCC GGG AGC TCA ACC CGG GTA AA TAC CCT

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F16.-13

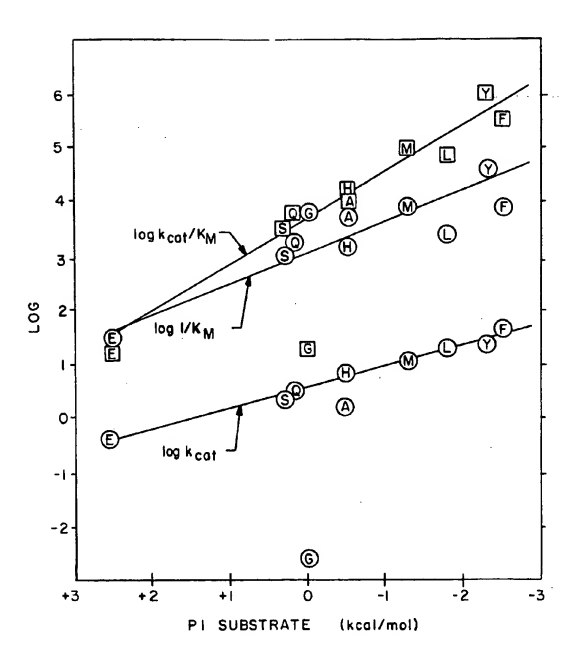


FIG. - 14

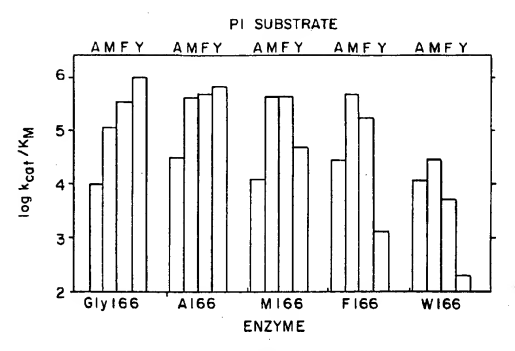


FIG. - 15A

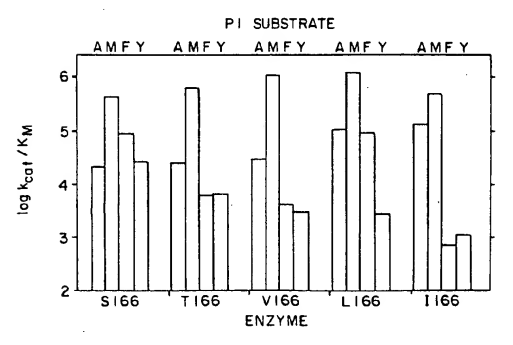
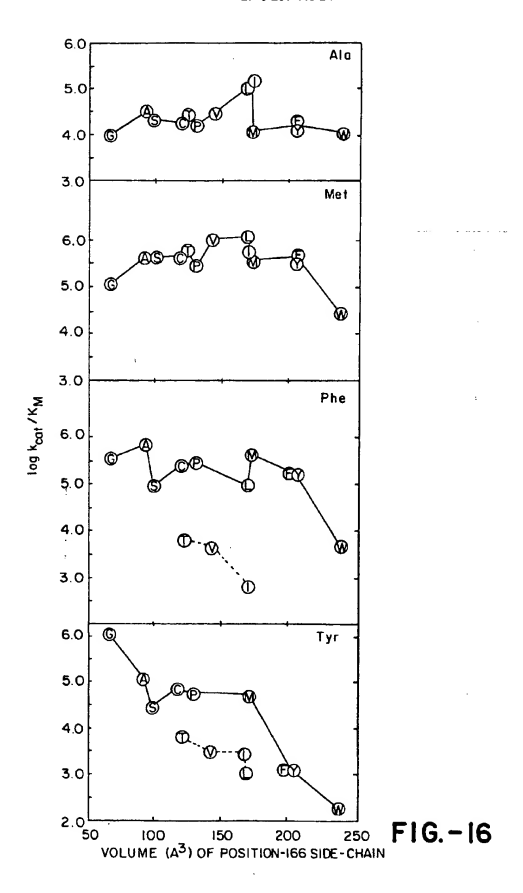
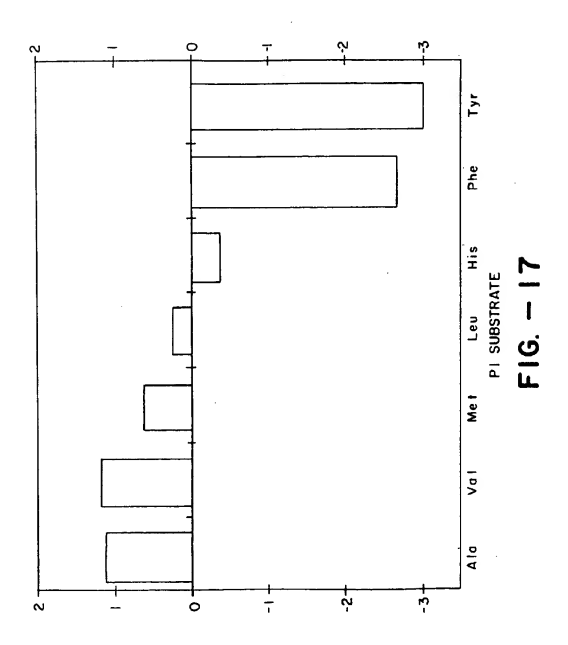


FIG.-15B

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GLY-169 CASSETTE MUTAGENESIS

Z	CODON: WILD TYPE AMING ACID SEQUENCE:	-	162 SER SER THR VAL GLY TYR PRO GLY LIS TYR PRO SER	ER TH	¥ *	ר פרי	7	PRO	169 GLY	. 517	Α.	17	₩ E	
	1. WILD TYPE DNA SEQUENCE	5	TCA AGC ACA GTG GGC TAC CCT GGT AAA TAC CCT TCT	SC AC	A GT	99 9	C TAC	. CCT	EGT	¥.	TAC (5	5	Ē
		'n	AGT TCG TGT CAC CCG ATG GGA CCA TTT	56 76	2	ני	3 ATG	66A	CCA	E	ATG GGA AGA	¥35	E	ŝ
										•	•			
2.	P169 DNA SEQUENCE		TCA AGC ACA GTC GGG JAC CCTGA TAL CCT TCT	SC AC	A GT	<u> </u>	IA	123		5	M	<u>[7</u>	ב	'n
		m	AGT TCG TGT CAC CCC ATG GGA KPNI	55 76	5	3	XPN K	66A		CT A	CT ATA GGA AGA ECORV	GA A	₩9	ŝ
											•			
m	P169 CUT MITH KPHI AND ECORVE	S	TAC AGC ACA GTC GGG TAC	SC AC	A GT	່ ເຄີຍ ເຄື່ອ	TAC				PAT CCT TCT	CT 1		'n
		ň	AGT TCG TGT CAC) 50 TG	T CA	ວິ					T ▲ 6	TA GGA AGA		5
						•					•			
÷	CUT P169 LIGATED WITH	S	TAC AGC ACA 6TG GGG TAC CCT NIN ANA TAT CCT TGT	SC AC	A GT	5 66	TAC	B	NNN	B	M	1	5	÷
	OLIGONUCLEOTIDE POOLS	3.	AGT TO	5 76	₹ -	3	AIG	TCG TGT CAC CCC ATG GGA NNN TIT ATA GGA AGA	NNN	日	TA G	SGA A		Š
Z	PLUTAGENESIS PRIMER FOR P169	2.5	AAG CAC AGT GGG GTA CCC TGA TAT CCT TCT GTC A	IC AG	1 666	5 67/	222 1	TGA	TAT	1 133	וכד פ	10 /		ň

5'-GGT-TCC-GGC-QAA-GCTT-AGC-TGG-ATC-ATT-3 5'-GGT-TCC-GGC-CAA-TAC-AGC-IGG-AIC-AIT-3' Gly-Ser-Gly-Gln-Tyr-Ser-Trp-Ile-Ile---TCC-GCC-CAA-NNN-AGC-TGG-ATC--104 105 2. Wild type amino acid sequence: 3. Wild type DNA sequence: 5. Primers for 104 mutants: 4. Primer for Hind III 1. Codon number: insertion at 104:

108

A,M, L,S, AND HI04

Mutants made:

5'-GTA-GTC-GTT-GCG-AGC-GCC-GGT-AAC-GAA-3' 5'-GTA-GTC-GTT-GCG-GCA-GCC-GGT-AAC-GAA-3' G-GTA-CCC-GGT-AAC-GAA-3' Val-Val-Val-Ala-Ala-Ala-Gly-Asn-Glu 5'-GTA-GTC-GTT-GC 2. Wild type amino acid sequence: 3. Wild type DNA sequence: 1. Codon number: 4. VI52/PI53 S 152: ĸ

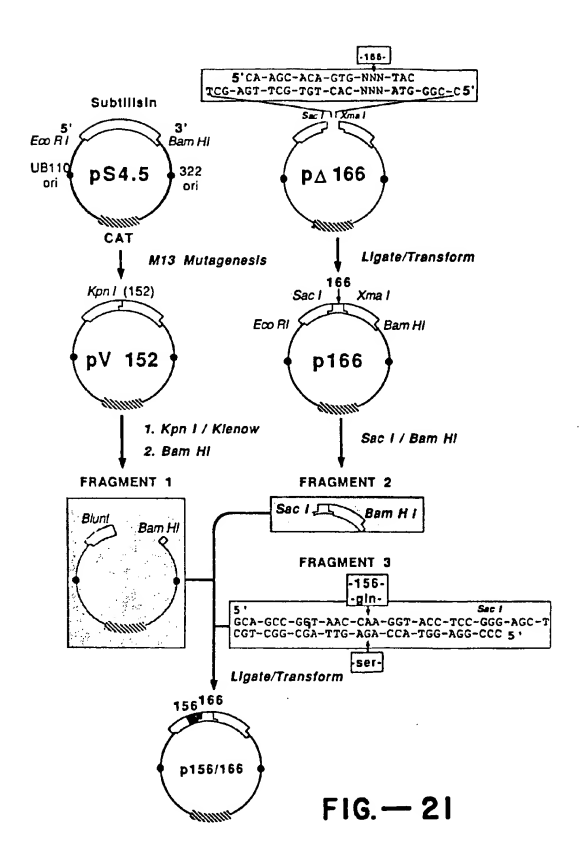
150

148

5'-GTA-GTC-GTT-GCG-GGC-GCC-GGT-AAC-GAA-3'

G 152:

ø.



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- ' \(\text{i} \) \(\text{i} \)	 Codon number: Wild type amino acid sequence: Wild type DNA sequence: 	1	215 -G1y-A1a- -GGG-GCG- -CCC-CGC-	217 Tyr-Asn-Gly TAC-AAC-GG1 ATG-TTG-CC	211 G1y-Asn-Lys-Tyr-Gly-Ala-Tyr-Asn-Gly-Thr-Ser-Met-Ala -GGA-AAC-AAA-TAC-GGG-GCG-TAC-AAC-GGT-ACG-TCA-ATG-GCA CCT-TTG-TTT-ATG-CCC-CGC-ATG-TTG-CCA-TGC-AGT-TAC-CGT-5.
4, <u>Q</u>	4. pó217	566A-AAC-AAA-IAC-66C-6CC-IAC66-AIA-IGA-AIG-6CA CCI-IIG-III-AIG-CCG-CGG-AIGCC-IAI-AGI-IAC-CGI Na'I	-990-990- -990-990-	180 6 6	GGA-AAC-AAA-IAC-GGC-IAC
ry g. g	5. pΔ217 cut with Nar I and Eco RI	5'-GGA-AAC-AAA-TAC-GG CCT-TTG-TTT-ATG-CCG-Gp	d9-922-		* pa-tca-atg-gca t-agt-tac-cgt-5'
ල <u>ර</u> ගු	6. Cut p∆217 ligated with cassettes:	5'-GGA-AAC-AAA-TAC-GGC-GCG-NNN-AAC-GGT-ACA-TCA-ATG-GCA CCT-TTG-TTT-ATG-CCG-CGC-NNN-TTG-CCA-TGT-AGT-TAC-CGT	-990-900- -909-299-	NNN-AAC-GGI NNN-TTG-CC	GGA-AAC-AAA-TAC-GGC-GCG-NNN-AAC-GGT-ACA-TCA-ATG-GCA CCT-TTG-TTT-ATG-CCG-CGC-NNN-TTG-CCA-TGT-AGT-TAC-CGT-5

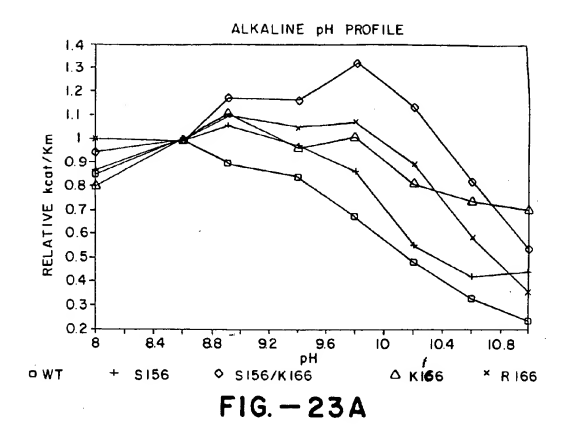
F16.-22

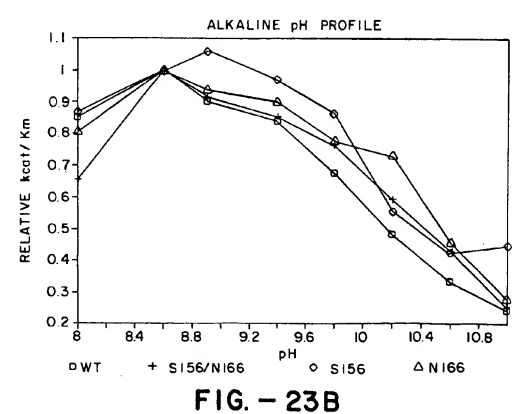
All 19 at 217

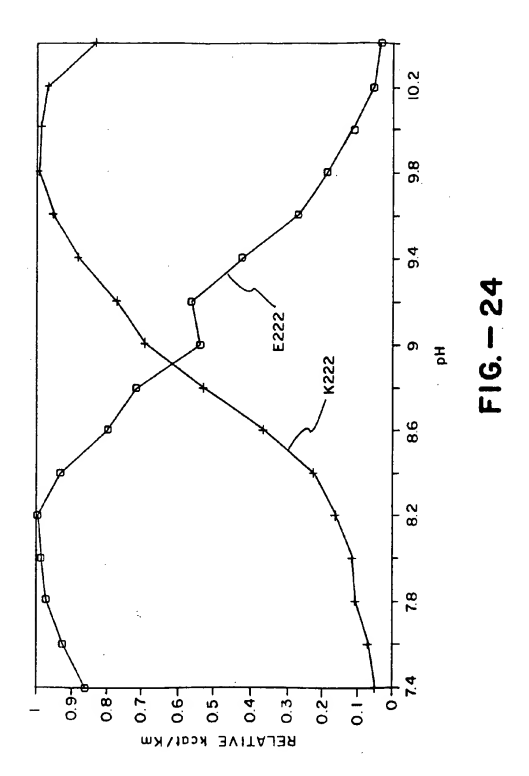
8. Mutants made:

5'-GA-AAC-AAA-TAC-GGC-GCC-TAC-GGA-TAT-CAA-TGG-CAT-3'

 Mutagenesis primer for p∆217;





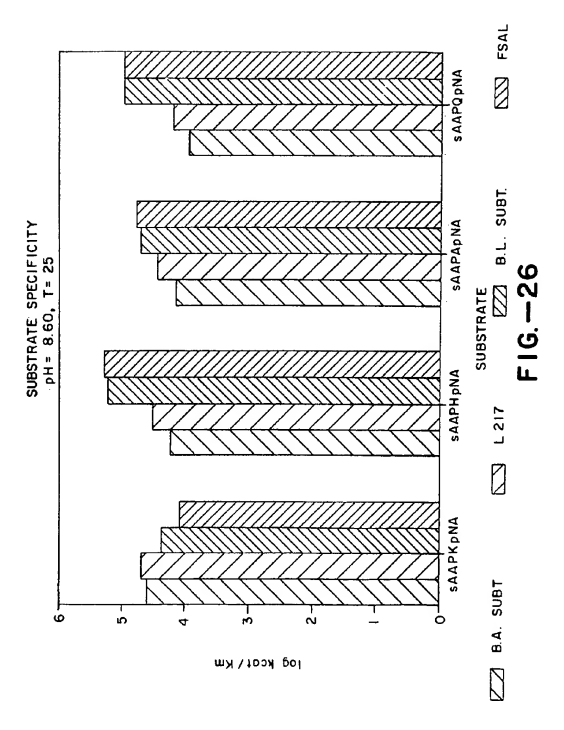


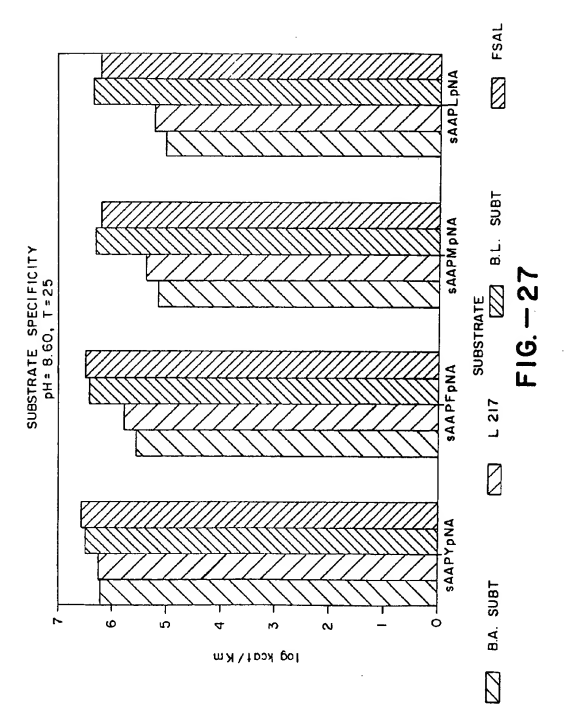
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4	 Codon number: Wild type amino acid sequence: Wild type DNA sequence: p∆95: 	Tyr-Ala-Val-Lys- 5'-TAC-GCT-GTA-AAA- ATG-CGA-CAT-TTT- 5'-TAC-GCG-T ATG-CGC-A	100 Tyr-Ala-Val-Lys-Val-Leu-Gly-Ala-Asp-Gly-Ser S'-TAC-GCT-GTA-AAA-GTT-CTC-GGT-GCT-GAC-GGT-TCC ATG-CGA-CAT-TTT-CAA-GAG-CCA-CGA-CTG-CCA-AGG-5' S'-TAC-GGG-TCTC-GGT-GCA-GGT-TCC ATG-CGC-AGAG-CGA-CGT-GCA-GGT-TCC	y-Ser T-TCC A-AGG-5' T-TCC
u,	5. pA95 cut with Muland Pst I	Mal 5'-TA * ATG-CGCp	* pGAC-GGT-TCC A-CGT-CTG-CCA-AGG-5'	T-TCC A-AGG-5'
9	6. Cut pA95 ligated with cassettes:	* 5'-TÂC-GCG-GTA-AAA- ATG-CGC-CAT-TTT-	* 5'-Tac-gcg-gta-aaa-gtt-ctc-ggt-gca-gac-ggt-tcc atg-cgc-cat-ttt-caa-gag-cca-cgt-ctg-cca-agg-5'	T-TCC A-AGG-5'
7	7. Mutagenesis primer for p∆95:	5'-CA-TCA-CTT-TAC-C	* * * * 5'-CA-TCA-CTT-TAC-GCG-T-CTC-GCT-GCA-GAC-GGT-TCC	GI-TCC

FIG. -25

8. Mutants made:





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